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Technoeconomic Evaluation of MEA versus Mixed Amines for CO₂ Removal at Near-Commercial Scale at Duke Energy Gibson 3 Plant

D. A. Jones, T. F. McVey, S. J. Friedmann

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Technoeconomic Evaluation of MEA versus Mixed Amines for CO₂ Removal at Near-Commercial Scale at Duke Energy Gibson 3 Plant

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Dale Jones, Tom McVey, and S. Julio Friedmann

Lawrence Livermore National Laboratory

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Executive Summary

Carbon emissions in the power sector remains an important goal for both US and Chinese decision makers and industrial leaders. In order to deeply reduce their greenhouse gas emissions, many utilities are considering carbon capture, utilization, and sequestration (CCUS). For the existing US and Chinese coal-burning fleets, post-combustion capture (PCC) is the only viable approach. While companies provide technology for PCC retrofits, typically solvent-based absorption systems involving amine solutions, the costs and performance of these systems remain largely untested in commercial settings. As such power generation companies have very few technical or economic data sets on which to base large capital decisions.

In support of gaining this information, the world's largest power company, Huaneng Power Group, has developed and deployed several PCC systems as retrofits to Chinese coal-fired power plants. Duke Energy Inc., North America's largest power generator, is considering options for CCUS retrofits and has created a partnership with Huaneng to assess the viability of a retrofit project on one of its US plants. In support of this effort, and in partnership with Duke and Huaneng under the US-China Clean Energy Research Program (CERC), LLNL has evaluated the economics of a novel amine-solvent technology for removal of CO₂ from flue gas developed by Huaneng's Clean Energy Research Institute (Huaneng CERI). This technology is being considered for implementation by Duke Energy at their Gibson-3 unit in Indiana.

Monoethanolamine (MEA), typically at 30% mass concentration, is a common technology used for amine solvent-based CO₂ capture. In this technology, CO₂ is absorbed into an amine-based solvent at low temperatures (<60°C) and then thermally regenerated at higher temperatures (100-130°C). A novel amine capture technology for CO₂ emissions has been developed by Huaneng Clean Energy Research Institute/Xi'an Thermal Power Research Institute, both part of Huaneng power. Huaneng's technology substitutes part of the MEA with hindered or tertiary amines, such as 2-amino-2-methyl propan-1-ol (AMP) or methyl diethanolamine (MDEA). These hindered amines have slower reaction kinetics than MEA, but somewhat better energetics of regeneration.

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We have concluded that Huaneng's technology represents an improvement, both energetically and economically over technology using solely MEA as the amine in the solvent. We assess it is capable of a 1.4% reduction in the generation loss over MEA alone (i.e. 19.7% generation loss as opposed to 21.1% loss). Also, we believe that the use of a solvent mix including amines that are less subject to degradation than MEA results in reduced solvent replacement costs. Our estimate is that use of the mixed amines could result in up to a 10% reduction in operating cost relative to 30% MEA. Huaneng is currently using its demonstration plants at Shidongkou and Gaobeidian to conduct tests to improve both the process configuration (for both solvent longevity and reduction of the energy penalty by heat integration). We further understand from Huaneng that further research on solvent composition is directed at extending the solvent life, rather than reduction in the energy penalty. Table ES.1 below summarizes our estimates of the costs of conventional MEA technology at Gibson-3 and the cost of our concept of Huaneng CERI's technology, at 30% and 35% total amine concentration.

Table ES.1. Cost Estimate Summary Table for 0.91 million tonne/annum CO₂ capture plant

Cost Parameter	30% MEA	Mixed Amines: 30%	Mixed Amines: 35%
Total Fixed Capital (US \$million)	\$133.4	\$135.8	\$128.9
Annual Operating Cost, including cost of Capital (US \$million)	\$59.0	\$57.4	\$54.6
Cost of CO ₂ Capture and Compression per tonne CO ₂	\$64.4	\$62.8	\$59.8
Estimated Impact on Power Generation Costs, \$/MWh	\$68.7	\$66.0	\$61.8

The results of this work will help US and Chinese companies better understand the likely cost and performance issues with PCC retrofits of coal plants. It also creates a baseline from which other technologies and potential projects can compare, hopefully leading to both emissions reductions and new commercial opportunities for US and Chinese companies.

Scope Note

At the request of the U.S. Department of Energy, we are evaluating a technology developed by China Huaneng Group for CO₂ capture. This technology has been proposed for use in a joint project between Duke Energy and Huaneng for a demonstration carbon capture and sequestration (CCS) process at unit 3 of Duke's Gibson Station in southern Indiana. Huaneng has modeled their process using Aspen Tech's software and verified its operation at smaller pilot plants at Gaobeidean and Shidongkou.

Our approach was to perform a technoeconomic analysis by creating independent process models. Based on the results of those models, we estimated the capital and operating costs of the process to determine the cost of generating electricity while capturing 90% of the carbon dioxide emitted by a fraction of the power unit.

The amine system we have chosen as our baseline is 30% monoethanolamine (MEA). Huaneng is developing proprietary novel amine systems that we approximated by a mixture, based on information in a Huaneng patent. We have presented the results for MEA at 30% concentration and the amine mixture at 30% and 35%.

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This report is presented in three sections, (1) Detailed description of the process chemistry and the results of the Aspen models for the three cases above, (2) Presentation of the process installation and operating costs for each case, and (3) Process modifications and improvements made necessary by the conditions at the Gibson Station and discussions with Huaneng engineers during a visit to the Shidongkou installation. Supplementary materials and tables are presented in appendices.

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Chapter 1: Engineering Analysis

Introduction

Beginning in June 2011, the LLNL, Huaneng, and Duke team began to exchange information in order to assess the merits of a PCC retrofit using the CERI technology. Early on, the decision was made to focus on a project at the Gibson power station in southwestern Indiana, in particular the number 3 unit. This selection was made based on several grounds, including the relatively high efficiency of the supercritical boiler there and the relative ease of engineering a retrofit in terms of space, cooling water, permitting, and other considerations.

Importantly, the three groups agreed to share information, data, and results for the purpose of objective analysis and assessment. In an unprecedented step, the parties also agreed to share operational information, site information, and cost data with the expectation of releasing this data. This arrangement and openness was critical to the success of the research program, and helps make the analysis unique in terms of its value as a potential benchmark for future retrofit technology assessments.

Process Review


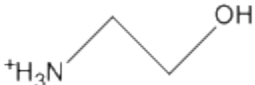
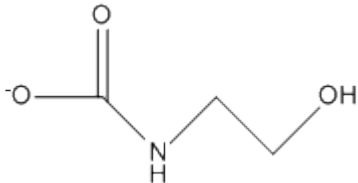
Discussion of Amine Technology

Alkanolamines, amines that contain an amine group and one or more hydroxyl groups, are a class of organic base that have been used to remove the acidic species, such as carbon dioxide from flue gas. These amines aid the absorption of CO₂ by either forming carbamate salts, or by stabilizing bicarbonate ions in solution. The carbon dioxide can then be removed as a gas by heating the CO₂-rich solution at moderate temperatures (90-130°C), after which the carbon dioxide gas can be compressed for storage or reuse. Alkanolamines (amines with an additional alcohol functional group) are generally used in the CCS process because of their increased solubility and reduced volatility compared to other amines.

Chemistry

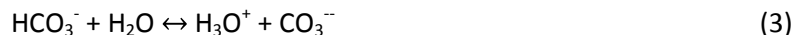
Discussion of chemistry of reactions

To evaluate Huaneng's new technology, we need to establish a baseline. We chose 30% aqueous monoethanolamine, as a solvent system that is used commonly both commercially and in modeling of CCS systems. Monoethanolamine forms two ionic species during the capture process, the amine cation, MEAH⁺, and the amine carbamate anion, MEACOO⁻, shown below.

		
Monoethanolamine (MEA)	MEAH ⁺	MEACOO ⁻

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The following equilibrium reactions take place in the process:¹



The CO_2 is, therefore, carried by the monethanolamine solution in the forms of bicarbonate, carbonate, and carbamate.

Not all amines used for CO_2 absorption can form carbamates. Tertiary amines such as methyldiethanolamine (MDEA) cannot form carbamates, and so-called “hindered” amines such as 2-amino-2-methyl-1-propanol (AMP) form carbamate with low stability. Such unstable carbamates as formed by AMP may play a significant role in kinetics of reaction, but do not act as significant carriers of CO_2 . These hindered and tertiary amines have a lower energy requirement for regeneration than MEA, but have slower kinetics, which can lead to higher overall solvent flow and larger equipment sizes. Components such as piperazine may be added to improve the kinetics of absorption. Therefore, optimization of the amine solvent composition is needed, and is non-trivial.

Huaneng Plants

Huaneng have three CO_2 capture plants working at present

- A pilot plant at Gaobeidian power plant, Beijing (3,000 tonnes/year CO_2)
- A newer pilot plant in Miyun county (capacity unknown)
- A demonstration plant at Shidongkou power plant, Shanghai (100,000 tonnes/year nominal, but debottlenecked to 120,000 tonnes/year)

The Shidongkou plant uses 30% MEA, but was commissioned using 20% MEA. The Miyun plant is used for testing of novel amine compositions. The Gaobeidian plant has used 20-30% MEA in demonstration runs, and may have also been used for testing of novel amine compositions.

Huaneng Patent

We based our assessment of the solvent composition used in Huaneng’s technology on a patent issued to Xi’an Thermal Power Research Institute, which lists several of the key personnel associated with the Huaneng amine- CO_2 project (e.g. Xu Shishen, Gao Shiwang, Liu Lianbo, amongst others) as inventors.² This patent describes compositions of amines to be used as CO_2 capture solvents, with the amine portion being between 50-85% MEA, 20-40% MDEA, and 0-10% of piperazine, with the balance of the amine being a sterically hindered amine (AMP, tertiary-butylaminoethanol [TBE], tertiary-butylaminoethoxyethanol [TBEE] or tertiary-butylaminopropoxyethanol [TBPE]). The patent describes the amines as being used in 10-40% aqueous solution, and with 0.02%-2% of sodium or potassium tetraoxovanadate. We believe, but cannot confirm, the tetraoxovanadate acts as a stabilizer either by inhibiting oxidative degradation or by catalyzing the oxidation of sulfite ions to sulfate.

Seven example compositions are given in the patent. We chose to model the example composition with the lowest proportion of MEA (50% MEA/40% MDEA/10% AMP), believing this would show the greatest difference in behavior from the base-case scenario using MEA as the sole amine. Informal conversations

with Huaneng have indicated that while Huaneng are still optimizing their solvent composition, the performance of what we modeled is reasonable for Huaneng's technology. They have also indicated that they have ruled out use of TBE, TBEE, or TBPE in their solvent mixture, as these are expensive relative to the other amines without offering a corresponding benefit for the increased cost.

Design Bases

Table 1.1 gives the design bases for the plant. Plant scale, on-line factor, flue gas temperature & humidity, and capture efficiency were specified by Duke Energy. Cooling water temperature was taken based on Duke Energy's description of peak ambient temperature. Thermal input and gross/net electrical output were estimated based on the fraction of flue gas from Gibson-3 needed to be treated to meet the required CO₂ capture rate. As noted above, the baseline 30% amine was selected by LLNL as being representative of commonly used amine technology, and mixed amine compositions were estimated based on the Huaneng patent. Flue gas compositions were based on data provided for Gibson-3 by Duke Energy.

Table 1.1. Design Bases

Specification	Value
Carbon capture rate	910,000 tonnes per year
Capture efficiency	90%
Plant on-line factor	80%
CO ₂ instantaneous input rate	1,262,500 tonnes per year
Fraction of Gibson 3 flue gas treated	25.5%
Thermal input	461 MW
Gross electrical output	177.6 MW
Net electrical output	161.9 MW
Flue gas temperature	55°C
Cooling water temperature	40°C
Amine System Case 1 - 30% by weight in water	MEA 100%
Amine System Case 2 - 30% by weight in water	MEA 50 wt%
	MDEA 40 wt%
	AMP 10 wt%
Amine System Case 3 - 35% by weight in water	MEA 50 wt%
	MDEA 40 wt%
	AMP 10 wt%

Flue Gas Composition at Gibson-3

The composition used in our designs for the flue gas for Gibson-3 is given in Table 1.2, based on data from Duke Energy. The composition is typical of that from a coal fired power plant. The carbon dioxide level of 9.51% is highly diluted with nitrogen as a result of burning the coal in air. The flue gas is saturated with water as it leaves the flue gas desulfurization (FGD) unit that reduces the sulfur dioxide (SO₂) level to 63 parts per million. Because SO₂ forms heat stable salts with amines, removal of these salts is necessary to prevent their buildup. The presence of oxygen in the flue gas can cause degradation of the amine. At the installation of interest, the flue gas leaves the FGD unit at 55°C, a temperature higher than the optimum temperature for absorption.

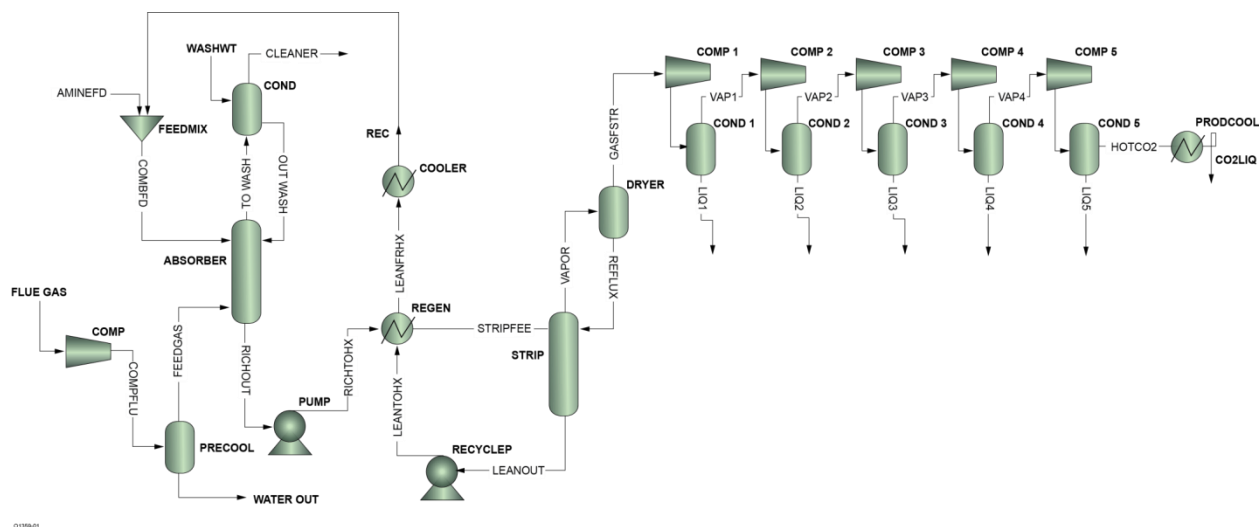
Table 1.2. Gibson-3 Flue Gas Composition

Component	Molar Composition,	Dry Gas Molar Composition,
Carbon dioxide	9.51%	11.25%
Water	15.51%	--
Sulfur dioxide	63 ppm	75 ppm
Nitrogen	67.62%	80.03%
Oxygen	6.55%	7.75%
Trace components (not included in ASPEN modeling)		
Hydrogen chloride	1.1 ppm	1.3 ppm
Ash	7.4 ppm	8.8 ppm
Nitric oxide (NO)	64 ppm	76 ppm
Nitrogen dioxide (NO ₂)	3.4 ppm	4.0 ppm
Argon	0.81%	0.95%
Carbon monoxide	17 ppm	20 ppm

Process Description

The process flow diagram is shown in Figure 1.1, with a larger version in Figure A.1 in Appendix A. We will discuss the process in sections 1) absorber and associated equipment, 2) stripper and associated equipment, and 3) the recycle loop. In the simulation of the three amine systems the unit operation configuration shown Figure 1 remained constant; however, since varying the amine compositions affected the efficiency of the carbon capture process, some equipment specifications and dimensions vary with the amine composition. Stream tables for the modeled processes are given in Tables B.1, B.2, and B.3 in Appendix B.

Figure 1.1. Carbon capture process



Absorber and associated equipment

The absorber (ABSORBER) is a packed column in which the cooled flue gas (FEEDGAS) from the precooler (PRECOOL) flows upward in contact with the lean amine solution (COMBFD) flowing downward. The carbon dioxide transfers from the gas phase to the liquid phase, to be converted to bicarbonate, carbonate, and carbamate ions as described in the chemistry section above. The cleaned flue gas (TO WASH) exiting the top of the column is submitted to a water wash and cooler (COND) to reduce amine emissions. For convenience

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in modeling we represent the wash and cooler as a separate item, but it actually is located in the top section of the absorber. The wash water (WASHWT) returns to the column (OUT WASH) and the effluent flue gas (CLEANER) is discharged to the atmosphere with 90% of its CO₂ removed.

Need for additional retrofit equipment: Absorption of CO₂ is thermodynamically more favorable at lower temperatures, and Huaneng used a flue gas feed temperature of 40°C in their model. Since the temperature of the flue gas from the FGD unit at Duke is 55°C, the flue gas must be precooled to give better absorption performance. Because it is saturated in water, cooling by water injection is not practical. The gas must, therefore, be cooled in a heat exchanger followed by a gas-liquid separator, (shown here as a single unit, PRECOOL), where excess water (WATER OUT) is removed. The cooling water available at the installation has a maximum temperature in the summer of 40°C, so, under such conditions, the absorber gas inlet temperature is at least 45°C. Because the influent gas to the CO₂ capture plant is near atmospheric pressure, a fan (COMP) provides the few inches of water increase in gas pressure to overcome the pressure drops in the precooler and the absorber column.

Stripper and associated equipment

The stripper (STRIP) is another packed column operated under conditions causing the carbon-containing species exiting the absorber to release gaseous CO₂, which can then be compressed and dried to 99.5% carbon dioxide. To reverse the equilibrium such that absorbed CO₂ can be released, the separator is operated at higher temperature than the absorber. The carbon dioxide rich solution from the absorber (RICHOUT) is pumped (PUMP) to a regenerative heat exchanger, described below (REGEN), and injected into the top of the stripper column, where it is stripped by an upflowing gas phase, created by the reboiler (not shown in the flow diagram) at the bottom of the column. This reboiler is the main source of energy loss in the carbon capture system. (The loss of energy can result in greater than a 20% reduction in the net electrical power of the installation.)

Need for additional retrofit equipment: At the top of the stripper is a condenser (DRYER) to reduce the water content of the CO₂ before it is sent to the compressors (COMP1 through COMP5). The separator is operated at about 1.5 atmospheres, to reduce the electrical requirement on the compressors, the second largest drain of the generated electricity. Our design includes a five-stage compressor with interstage cooling (COND1 to COND5) to produce a compressed product at 153 bar (CO2LIQ), containing over 99.5% CO₂.

The stripper reboiler requires careful temperature control. At Gibson-3, the low pressure steam supply is at approximately 11 bar and 330°C, depending on the quantity of steam extracted from the LP steam turbine: however the LP steam pressure varies according to the operating conditions in the power plant. To prevent thermal degradation of the amine at the reboiler heat exchange surface, the maximum temperature of reboiler steam should not exceed ~120°C, corresponding to 2 bar pressure for saturated steam. To achieve this, the 11 bar steam is expanded through a letdown turbine, which generates electricity to partially offset the power required to compress the product CO₂. Because the use of saturated steam results in a higher heat transfer coefficients and smaller reboiler, water is added to the steam supplied to the stripper reboiler to take the steam from a superheated condition to close to saturation.

A structured packing is used in both the absorber and stripper columns. Structured packing, although typically more expensive than random packings, gives better contact area per unit volume, reduced risk of flooding from excessive gas flows, and reduced pressure drop across the column, relative to a random packing.

Recycle loop

The purpose of the recycle loop is to heat the CO₂-rich solution (RICHTOHX) for injection into the stripper while cooling the CO₂-lean solution from the stripper (LEANTOHX) for injection into the absorber. The main component of the recycle loop is the regenerative heat exchanger (REGEN) that heats the stripper feed (STRIPFEE) as it cools the stripper lean (LEANFRHX) solution. The recycle loop contains two centrifugal pumps, one (PUMP) to advance the CO₂ rich absorber product through the regenerative heat exchanger and compress it to 1.5 bar as it is fed to the stripper. The recycle pump (RECYCLEP) overcomes the pressure drop in the regenerative heat exchanger, and the head difference between the bottom of the stripper and the top of the absorber column. The cooler (COOL), reduces the recycled solution temperature to the desired absorber feed temperature. The recycled solvent is adjusted for amine and water concentration (makeup solvent is the AMINEFD stream) before being fed to the absorber. The stream, AMINEFD, in Tables B.1 – B.3 contains a significant amount of water and a very small (MEA case) to not measurable (Mixed amine case) amount of amine. This stream also includes the water added to the washer (COND) at the top of the absorber.

Baseline 1: CO₂ Removal using 30% Monoethanolamine

Process Details

Major equipment items are listed in Table 1.3. A stream table is given in Table B.2 in Appendix B.

Table 1.3. Baseline Case – Solvent Composition of 30% MEA Major Process Equipment List

Item Label in Process Flow Diagram (PFD)	Number	Size	Description	Comments
Columns				
ABSORBER	2	25 m ht x 9 m dia	304 SS column shell 15 m of 316 SS structured packing	Vapor wash step at top of column to reduce amine emissions is modeled as COND
STRIPPER	2	10 m ht x 4.8 m dia	304 SS column 6 m of 316 SS Structured packing	
Pressure Vessels				
DRYER	2	7 m x 3.5 m dia	Knock-out drum for stripper	304 SS
COND1	1	5 m x 2.5 m dia	First CO ₂ compression stage knock-out drum	304 SS
COND2	1	3.0 x 1.5 m dia	Second CO ₂ compression stage knock-out drum	304 SS
COND3	1	1.8 x 0.9 m dia	Third CO ₂ compression stage knock-out drum	304 SS
COND4	1	1.2 x 0.6 m dia	Fourth CO ₂ compression stage knock-out drum	CS
COND5	1	0.6 x 0.3 m dia	Fifth CO ₂ compression stage knock-out drum	CS
Heat Exchangers				
REGEN	14	1,000 m ²	660 GJ/hr exchange	Shell: 304 SS Tubes: 304 SS
DRYER	2	1,000 m ²	-115 GJ/hr cooling	Shell: CS Tubes: 304 SS

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Item Label in Process Flow Diagram (PFD)	Number	Size	Description	Comments
REBOIL	4	1,000 m ²	440 GJ/hr heating	Shell: CS Tubes: 304 SS
COOLER	4	1,000 m ²	-137 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND1	4	900 m ²	-18 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND2	3	800 m ²	-12 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND3	3	800 m ²	-12 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND4	3	800 m ²	-12 GJ/hr cooling	Shell: CS Tubes: CS
CO2COND5	4	1,000 m ²	-24 GJ/hr cooling	Shell: CS Tubes: 304 SS
PRECOOLERHX	10	1,000 m ²	-108 GJ/hr	Shell: CS Tubes: 304 SS Not shown on flow diagram
INTERCOOLER	2	1,000 m ²		Shell: CS Tubes: 304 SS Not shown on flow diagram
Compressors				
COMP	7	0.1 MW	Pre-Absorber Blower	Stainless Steel
COMP1	1	2.25 MW	Brake Power	Stainless Steel
COMP2	1	2.25 MW	Brake Power	Stainless Steel
COMP3	1	2.25 MW	Brake Power	Stainless Steel
COMP4	1	2.25 MW	Brake Power	CS
COMP5	1	2.25 MW	Brake Power	CS
Pumps				
Absorber Bottoms Pump	5	0.3 m ³ /s	750 kPa ΔP	304 SS
Stripper Bottoms Pump	5	0.3 m ³ /s	300 kPa ΔP	304 SS
Intercooling Pump	5	0.3 m ³ /s	30 kPa ΔP	304 SS. Not shown on flow diagram
Tankage				
Amine Solution Surge Tank	2	750 m ³		304 SS
Amine Solution Mixing Tank	1	90 m ³		304 SS
Caustic Mixing Tank	1	10 m ³		Glass-lined CS.
Caustic Storage Tank	1	400 m ³		Glass-lined CS. 28 day storage
MEA Storage Tank	1	180 m ³		304 SS. 28 day storage
Other				
Ion Exchange Column	2	7.75 m x 2.5 m dia	Used to remove heat-stable salts from the amine solution	Resin-lined CS pressure vessel. Not shown on flow diagram
Activated Carbon Beds for Amine Solution	2	6 m x 2 m dia	Used to remove amine degradation byproducts from amine solution	Activated carbon bed with resin-lined CS. Not shown on flow diagram

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Item Label in Process Flow Diagram (PFD)	Number	Size	Description	Comments
CO ₂ Drying Beds	2	5.5 m x 1.5 m dia	Used to dry CO ₂ after COND3 knock-out drum	Silica: CS vessel shell. Not shown on flow diagram
Activated Carbon Beds for treatment of CO ₂	2	4 m x 1 m dia	For removal of organic impurities from CO ₂ .	Activated carbon with resin-lined CS. Not shown on flow diagram

MEA – 30% Amine Concentration

A process flowsheet showing most, but not all, of these equipment items is given in Appendix A as Figure A.1. A stream table of process flows for the baseline 30% MEA scenario is given in Table B.1.

Mixed Amines – 30% Amine Concentration

The stream table for the 30% Mixed amine scenario is given in Table B.2 in Appendix B; the corresponding equipment list is given in Table C.1 in Appendix C. The sizes of major equipment items change only slightly compared to 30% MEA base-case scenario. There is some additional tankage related to feed amine storage and mixing compared to the base-case, due to the use of three different amines rather than only one.

Mixed Amines – 35% Amine Concentration

The stream table and equipment list for 30% mixed amines is given in Table B.3 in Appendix B; the corresponding equipment list is given in Table C.2 in Appendix C. The sizes of major equipment items change only slightly compared to 30% MEA base-case scenario, but are slightly smaller than the 30% mixed amine scenario. Similar to the 30% mixed amine scenario, there is some additional tankage related to feed amine storage and mixing compared to the base-case.

Power Losses

The power loss values presented in Table 1.4 are to be compared with the output of 25.5% of the output of the Duke Energy Gibson-3 unit or 461.3 megawatts thermal (MWTh) with a gross generation level of 177.6 megawatts electrical (MWe) and a net electrical output after peripheral equipment losses of 161.9 MWe. The three amine systems compared in this table are the same as described above.

Table 1.4. Table of Comparative Power Losses for all three processes

	30% MEA	30% Amine Mix	35% Amine Mix
Reboiler MWTh used	122	115	109
Reboiler GJ/MT*	3.6	3.16	3.02
Electrical MWe used	11.9	11.9	11.8
Turbine MW generated	9.6	9.2	8.9
Cooling MWTh	127	135	126
LP steam diverted, kg/h	192,000	181,000	172,000
Percentage energy loss based on thermal input and electrical output			
Based on electrical output of 161.9 MWe	21.1%	20.3%	19.7%

*GJ = gigajoules

MT = metric ton of recovered CO₂, tonne

Materials of Construction

Because of the corrosive nature of carbon dioxide with water, the assumed material for equipment in contact with the recirculating solvent was predominantly 304 stainless steel. The packing in the columns is 316 stainless. Activated carbon tends to corrode stainless steel; therefore the beds using ion exchange resin or activated carbon were assumed to be resin- or polymer-lined steel. Vessels containing caustic were assumed to be glass-lined steel. In the product CO₂ compression train, it was assumed that CO₂ would be dried using a silica bed after the third compression/cooling step, permitting the fourth and fifth compression stages to use a less corrosion-resistant grade of steel.

Removal of Waste Products from the Solvent

Specific removal and treatment of trace contaminants was not included in our modeling using ASPEN. However, equipment we included in our cost estimates addresses removal of some of the contaminants. For example, sulfur oxides, nitrogen oxides, and hydrogen chloride lead to the formation of heat-stable salts with the amine species in the solvent solution. We have included in our capital and operating costs the cost of ion exchange treatment to remove such heat stable salts. Additionally, we have included activated carbon beds for removal of some of the products from amine degradation. We have sized this equipment based on an assumed 10% purge stream from the amine recycle.

At the Shidongkou plant, Huaneng uses vacuum distillation to recover MEA solution from heat-stable salts. However, we do not believe such a process would be optimal for mixture of MEA/AMP/MDEA, as the lower vapor pressures of AMP and MDEA would prevent effective recovery of those compounds: for instance, the vapor pressure of MDEA is 10 mmHg at 128 C.³ We judge this low vapor pressure would preclude recovery of the mixed amine by vacuum distillation, as temperatures above 120°C result in increased thermal degradation of MEA.

Additional mechanical filtration of solvent prior to ion exchange and activated carbon beds may be needed to avoid blinding of such beds by ash particles from the flue gas captured during absorption. We have not included such filters in our cost estimate, but judge such filtration systems for the purge stream would be minor compared to other cost items.

We note, based on information from Duke Energy, that selenium accumulation has caused waste solvent to have to be disposed of as a hazardous waste in other pilot tests of CO₂ capture using amines. Selenium is challenging to remove below RCRA limits by conventional methods, because of its chemical similarity to sulfur. We have assumed in our cost estimates that waste solvent disposed is concentrated ~50% by evaporation before disposal as hazardous waste.

Other Waste Streams

Other waste streams to be disposed of, as well as spent solvent, include brine from regeneration of the ion exchange columns, spent carbon from the activated carbon beds, and spent ion exchange resin. Estimated costs for treatment or disposal of these wastes have been included in the cost estimates presented later in this report. (See Chapter 3)

Chapter 2: Process Economics

Summary

The likely configuration, design, and operational requirements of a Huaneng-CERI based retrofit project on Gibson provided enough information to assess the likely capital and operational costs of the project. Tables 2.1 and 2.2 summarize the estimated process economics for each scenario. Full tables giving the variable, capital, and operating costs for the three scenarios analyzed are given in Appendix D.

Table 2.1. Capital Cost Summary

Capital Costs, US \$million (2011)	30% MEA	30% Mixed Amines	35% Mixed Amines
Battery Limits Investment	\$97.1	\$99.1	\$93.4
Offsites and Utility Investment	\$37.2	\$36.7	\$35.5
Total Fixed Capital	\$134.4	\$135.8	\$128.9

Capital costs for the baseline 30% MEA scenario are given in Table D.2 in Appendix D. Capital costs for the 30% and 35% mixed amine scenarios are given in Table D.5 and D.8, respectively, in the same appendix.

Table 2.2. Operating Cost Summary

Operating Costs, US \$million (2011)	30% MEA	30% Mixed Amines	35% Mixed Amines
Variable Costs			
Materials Consumed and Waste Disposal	\$5.8	\$4.3	\$4.3
Utilities	\$3.6	\$3.7	\$3.5
Lost Power Generation	\$14.1	\$13.6	\$12.9
Total Variable Costs	\$23.5	\$21.6	\$20.7
Fixed Costs			
Direct Labor	\$3.3	\$3.4	\$3.2
Maintenance Labor and Supplies	\$3.0	\$3.0	\$2.8
Plant Overhead, Taxes, and Insurance	\$4.8	\$4.8	\$4.6
Levelized Depreciation	\$4.4	\$4.5	\$4.3
Corporate Overhead	\$2.0	\$1.9	\$1.8
Cost of Capital	\$17.9	\$18.1	\$17.2
Total Operating Costs	\$59.0	\$57.4	\$54.6
Cost per tonne CO ₂ captured, US\$/tonne	\$64.4	\$62.8	\$59.8
Additional cost of CO ₂ captured per MWh generated, US\$/MWh	\$68.7	\$66.0	\$61.8
Reduction in costs compared to base case 30% MEA	--	4%	10%

CO₂ plant capture capacity is 1 million short tons (0.91 million tonnes), with 90% capture efficiency.

Operating costs for the baseline 30% MEA scenario are given in Table D.3 in Appendix D. Capital costs for the 30% and 35% mixed amine scenarios are given in Table D.6 and D.9, respectively. Variable cost estimates are given in Tables D.1, D.4, and D.7 in Appendix D for 30% MEA, 30% mixed amines, and 35% mixed amines, respectively.

Conclusions and Analysis

We assess that use of the mixed amines, as envisioned in the Huaneng patent, can give modest but definite benefits: up to 10% in operating cost savings for 35% mixed amines. The savings primarily come from both lower lost generation and from reduced material costs because of increased amine stability.

We conclude that the Huaneng mixed amine technology represents an improvement over conventional 30% MEA, but that the improvement in operating cost is not only due to reduced heat of regeneration, but also due to lower amine losses from degradation reducing the solvent replacement costs. We understand that Huaneng is working on process improvements to both decrease lean loading (and hence reduce solvent flow), to improve heat integration, and further extend solvent life. We envision these could yield an additional modest improvement in economics when implemented. These improvements are described in Chapter 3.

Methodology

A factorial method based on equipment costs was used for capital cost estimation. Freight-on-board (FOB) costs of capital equipment were estimated based on the equipment size, using published correlations for equipment costs, and adjusted for materials of construction, and inflated to 2012 costs. Installation costs for the equipment were estimated as factor (multiple) of the FOB equipment costs. Additional adjustments were made to account for the difficult geotechnical conditions at the site, and the fact that the installation would be a retrofit rather than a greenfield installation.

After capital costs were estimated, variable costs (including the chemicals consumed, electricity generation losses, and utility costs) were generated based on the mass and energy balance. From the capital and variable costs, operating costs were estimated.

The variable components of the operating costs are costs that are expected to scale with the quantity of CO₂ absorbed. These costs include not only the losses to electricity generation, but also chemical costs from operation of the CO₂ capture plant (predominantly solvent makeup costs).

Additional operating costs were added for operating labor and plant labor overhead; maintenance labor and supplies; property taxes and insurance; depreciation; cost of capital; and corporate-level general & administrative charges. Depreciation was treated as straight-line over thirty years.

Equipment Sizing

Major equipment items were sized based on either simulation results or on standard rules of thumb for conceptual design of chemical plants. Column diameters were calculated based on ASPEN simulations. Heat exchanger areas were calculated based on using the log-mean temperature difference and typical values of heat transfer coefficients based on nature of the fluids on either side of the heat transfer surface: shell and tube heat exchangers were assumed. Compressors and fans were sized based on their power rating; pumps were sized based on pressure difference and flow. Process vessels were sized based on flow rates through said vessels and required hold-up times. Tankage volume was calculated based on the flow rates and the assessed necessary hold-up time (for surge vessels and mix tanks). For feed storage tanks, we assumed the tanks held 2-4 week inventory of chemicals. Ion exchange columns were sized assuming a 1 hour empty bed contact time. Activated carbon beds were sized assuming an empty bed contact time of 0.5 hours.

Capital Cost Estimation

Freight-on-board cost of capital equipment items were estimated using correlations in Peters & Timmerhaus, 5th Edition, 2002, or using specific vendor quotes.⁴ Costs were inflated to 2011 prices using

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the Chemical Engineering (a trade magazine) Cost Index (CECI). Capital costs were separated into Battery Limits Investment (BLI) including equipment cost and installation of process equipment handling process streams, and Outside Battery Limits Investment (OBLI) which includes utilities, tankage, and general service facilities, and pipelines for utilities and flue gas from the main Gibson plant. Utility investment includes construction of cooling water towers; steam and any make-up process water were assumed to be supplied by the main Gibson plant

Installation costs for equipment were estimated using a factorial method. Installation costs were broken down into four components:

- Construction and fabrication (see discussion below)
- Instrumentation (15% of base equipment costs)
- Electrical (30% of base equipment costs)
- Piping (60-90% of base equipment costs)

Base equipment cost is the freight-on-board cost for a piece of equipment fabricated in carbon steel. A material factor of 2.2 was used for equipment assessed to be 304 stainless steel compared to the carbon steel cost. For equipment assessed to be stainless steel, piping costs were assumed to increase but not construction, electrical, or instrumentation costs.

Special Considerations for the Gibson Site

Geotechnical conditions are difficult at the site. The only area available for construction would be part of the current ash disposal area. Further, the underlying soil in the area is poorly consolidated, and the area is seismically sensitive. Hence, it was assumed that the construction cost component of installation costs would be raised by a factor of 2.5 to reflect additional civil engineering costs, such as driving piles to the approximate 50' bedrock depth. Hence, for carbon steel equipment the construction and fabrication costs were assumed to be between 150%-225% of the freight-on-board equipment costs. For 304 stainless steel, the construction and fabrication costs varied from 68%-102% of the equipment cost.

As costs in Peters & Timmerhaus are for greenfield construction, a complexity factor of 1.2 was used to reflect the additional design engineering and construction costs of constructing a facility within an existing operating plant.

Comparison with Capital Costs for Shidongkou

We assess, by comparing to the cost of the Shidongkou plant, that our capital cost estimates are in the reasonable range. A method of cross-checking chemical plant capital cost estimates is to use construction costs for previously constructed plants, and adjust for costs related to scale and location. The Shidongkou plant was installed at a cost of RMB 160 million (US\$24 million) for a nominal capacity of 100,000 tonnes using 15-20% MEA, later debottlenecked to 120,000 tonnes. Capital cost for chemical plants built in China vary between 60-80% of the cost for an equivalent U.S. plant. Therefore, using a location factor of 70% to convert from U.S. costs to Chinese costs⁵, and a scaling exponent of 0.66, we would get rough estimate of a capital cost for US\$130 million for a 910,000 tonne capacity plant in the U.S. However, the Shidongkou plant uses a 15-20% solution of MEA, so would have a somewhat higher capital cost than a plant using 30% MEA. However, this is balanced out by the need in our cost estimates to account for poor geotechnical conditions at the Gibson power station. Given this, our estimate of the capital cost of an MEA carbon capture plant of US\$134 million seem to be in line with what we would expect from comparison with the plant at Shidongkou.

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Operating Costs

Variable costs for the processes include materials consumed, the opportunity cost of electricity not generated because of operation of the plant, and utilities used (cooling water). Steam and any make-up process water were assumed to be supplied by the Gibson plant; the cost of supplied steam is costed as opportunity loss of lost electricity generated.

Cost of amines consumed were based on list prices given by Dow Chemical, which were then discounted by 20%, as contract prices for chemical are typically substantially lower than list prices. Costs of other materials (e.g. activated carbon, ion exchange resins, desiccants) were based on professional experience and data found on vendor websites on prevailing prices. Estimates for amine were based on MEA degradation rates using data from Singh, et al.;⁶ relative degradation rates for MEA and MDEA relative to MEA were extrapolated from published experiments by Lepaumier et al. for aqueous amines at 140 C and high pressures of O₂ and CO₂,⁷ and also based on feedback from Huaneng.

While the carbon capture plant is operating, there is a loss of electricity generation due to both thermal energy diverted to the carbon capture plant for solvent regeneration, and also due to parasitic electricity demand of pumps and compressors used in the process (including the cooling water recirculation pumps). The cost of the energy penalty of steam diverted from production of electricity to the stripper reboiler was priced as an opportunity cost, using the five-year average contract price for wholesale electricity for Midwest ISO/Cinergy, which is the grid Independent System Operator for the Gibson plant

Cost Parameters

Table 2.3 indicates the cost parameters used by the team in the capital and operating cost estimates.

Table 2.3. Cost Parameters Used

Cost Parameter	Value Used	Comments
Location	Princeton, Indiana, USA	Project-Specified
Discount Rate	12.5%	Team Estimate based on typical energy firm cost of capital
Depreciation Lifetime	30 years	Industry Standard; treated as straight-line (levelized) depreciation
Contingency factor for capital costs and for drilling costs	20%	Team Estimate
General Service Facilities	20% of Total Fixed Capital	Team Estimate
Waste Treatment Facilities	5% of Battery Limits Investment	Team, Estimate
Labor costs	\$62/hour	Bureau of Labor Statistics Manufacturing Compensation in Indiana
Plant overhead	80% of Operating Labor	
Maintenance Costs	6% of BLI	Typical level for high-maintenance chemical processes
Taxes & Insurance	1.6% of BLI	Industry Standard
General, Admin, Sales & Research	5% of Plant Gate Costs*	Team Estimate

* Plant gate costs are defined here as the cash cost plus depreciation charges. Production cost is equal to the plant gate cost plus a charge for corporate general, sales, administration and R&D costs (GASR). Operating costs are defined as the production cost plus the cost of capital (i.e. the capital investment times the discount rate).

Implications for retrofit economics

As discussed in the conclusions subsection above, we estimate that the Huaneng process, especially at 35% concentration, shows modest but definite cost benefits over 30% MEA: with our assessment of the Huaneng process showing up to a 10% operating cost advantage. Capital cost advantages are only seen when using 35% mixed amines, and are slight (3%); however the advantages in energy consumption and solvent degradation are more substantive and are seen for both 30% and 35% mixed amine concentration. We estimate that the cost of CO₂ capture is approximately \$60/MWh. Note this does not include the cost of liquefaction and disposal of the CO₂ captured from the plant, although it includes the cost of compression of CO₂ to 150 bar.

Chapter 3: Uncertainties and Opportunities

Uncertainties

As noted in previous chapters, since we did not have access to the rate-based absorption/distillation module in ASPEN, we had to approximate the column behavior by using equilibrium models for the absorber and stripper. This introduces uncertainty into the packed height of these columns (as we used a published correlation to estimate the HETP for the columns⁸, rather than being able to use rigorous simulation), and also leads to a potential underestimation of the total solvent flow. However, given the availability of components for accelerating the rate of absorption (e.g. piperazine), we believe we are likely within 10% of the total solvent flow.

While we have some remaining uncertainty regarding the energy loss due to the stripper reboiler in the mixed amine case, we have been informed by Huaneng that they do not anticipate being able to reduce the energy loss to the stripper reboiler much below 3.0 GJ/tonne CO₂ absorbed: there may be some reductions from better heat integration, but they anticipate their future focus on solvent composition will be on extending solvent lifetime rather than on reducing the energy penalty.

We are uncertain about the necessary amount of process water make-up, because changes in the precooling operation and in the handling of water in the solvent purge stream sent to purification will radically affect the process water balance. However, even if the most conservative estimate of the water imbalance (the water content in the AMINEFD stream) was made up using process water, the impact on operating cost would be less than \$150,000/year; this is minor compared to the overall variable cost estimates of ~\$20 million/year. Further, we do not believe any of the three scenarios considered would have a substantive advantage in process water consumption over another.

Opportunities for Further Optimization

We understand that Huaneng is pursuing several strategies for further heat integration and for extending solvent life. These include:

1. Intercooling in the absorber
2. Multiple feed locations of lean solvent to absorber
3. Bleed, alkalination, flash and mechanical vapor recompression (MVR) of the lean solution
4. Cooling of incoming raw flue gas with rich solvent
5. Multiple flashes of rich feed to stripper

Detailed description of these opportunities follows:

1. **Intercooling in the absorber.**

As the amine solvent absorbs CO₂, the heat of reaction of MEA (and other amines) with CO₂ leads to heating of the solution. As higher temperatures shift the equilibrium constants of the solvent values to those less favorable for absorption, cooling of the solvent during absorption reduces temperature peaks in the absorber and decreases solvent flow. A typical intercooling design might have 1-2 intercooling stages between packing sections. Solvent is pumped out of the absorber, cooling in heat exchangers, and then pumped back into the column into the distributors between packed sections.

Some intercooling is included LLNL's model. We found that intercooling is more effective for solvents with a higher heat of absorption of CO₂ (i.e. while intercooling improves the absorber performance for all solvent mixtures, the improvement was greater for 30% MEA than for the mixed amines). Intercooling of the liquid in the absorber can be used to eliminate the precooling

of the flue gas from 55°C to 44.4°C at a lower capital cost because liquid cooling requires a smaller heat exchanger than gas cooling for the same quantity of heat transferred. A disadvantage of using intercooling to replace precooling is that the 44 tonnes per hour of water that the precooler removes would then be added to the circulation system, requiring water bleed and amine loss.

2. Multiple Feed locations of lean solvent into the absorber.

Huaneng expect that by introducing the lean CO₂ at multiple locations in the absorber, they can reduce the partial pressure of CO₂ in the upper part of the absorber column, by causing more of the absorption of the CO₂ to occur in the lower sections of the absorber. With this reconfiguration, they expect that the lean CO₂ fed to the upper reaches of the column will further reduce the partial pressure of CO₂. As with intercooling, the aim is to reduce the overall solvent flow while maintaining the same performance of CO₂ removal from the flue gas.

3. Bleed, alkalination, flash and mechanical vapor recompression (MVR) of the lean solution.

This more unconventional system adds a bleed stream and pressure swing step to the CO₂ stripping section. In this configuration, a slip stream of 10% of the lean solution is bled from the stripper bottoms discharge and mixed with sodium hydroxide to convert heat stable amine bisulfite and sulfite salts to sodium bisulfite and sulfite and free amines. The resulting mixture is flashed to below-atmospheric pressure to volatilize CO₂, steam, and amine, leaving heat stable sodium bisulfite (hydrogen sulfite) and sulfite salts in the residual liquid. The vapor is recompressed and fed to the stripper bottoms section. The aim of this configuration is to lower the lean loading of CO₂ on the solvent, both by flashing some CO₂ in the flash drum, but also, as most of the recompressed vapor is steam, the partial pressure of CO₂ in the lower part of the stripping column is reduced, further reducing the lean loading. For the mixed amine scenario, as a portion of the low boiling monoethanolamine (MEA) is replaced with methyldiethanolamine (MDEA) and 2-amino-2-methylpropanolamine (AMP), the amine boiling point will increase and amine recovery in this step will be more difficult than for the scenario where solely MEA is used.

4. Cooling of incoming raw flue gas with rich solvent

This is a more straightforward heat integration step, using low-grade heat from the flue gas prior to the flue gas desulfurization to heat the rich solvent before entering the stripping column. This should lower the reboiler duty. Disadvantages are that this type of heat integration can be difficult if the pipe runs are long, and the poor gas-side heat exchange coefficients can result in large heat exchanger sizes for only a moderate amount of heat integration. Also, as the proportion of CO₂ captured from the plant increases, the increase in temperature of the solvent from pre-heating by the flue gas decreases, i.e. as the CO₂ capture process scales up, the impact of this process change per tonne of CO₂ captured will decrease.

5. Multiple flashes of rich solvent prior to feed into the stripper.

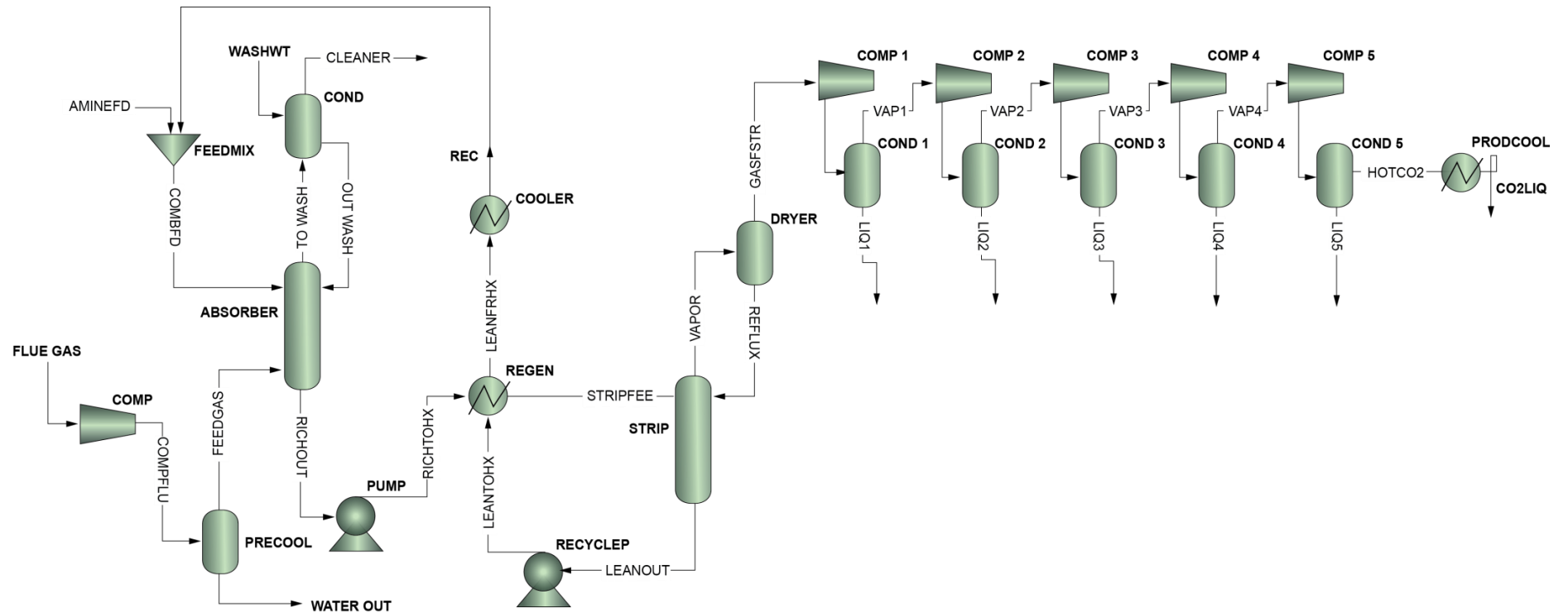
Unlike the flash stages after the stripper, here the aim is to reduce oxidative degradation of the amine in the stripper by flashing off most of the oxygen prior to the solvent being exposed to the higher temperature in the reboiler.

Additional implications for potential operations

Huaneng has been able to directly assess process improvements at their three pilot-to-near production scale amine-based carbon capture facilities in China. Although we have not performed formal modeling of the process improvements mentioned above, based on our qualitative assessment we see them as incrementally improving the operability and economics of a carbon capture plant. For instance, although a pre-stripper flash will decrease oxidative degradation of the amines, there are still thermal- and CO₂-catalyzed degradation paths that would lead to solvent degradation, particularly for MEA. While the cost of solvent replacement would decrease, we qualitatively assess this would be at most a 20-30% decrease in solvent replacement costs. Further, the process improvements, while reducing the energy penalty (and hence cost of lost electricity generated) and the costs from replacement solvent, would not be expected to radically reduce capital-related and other fixed costs, which are over 60% of the total costs. Hence, these improvements may add further to the modest operational cost benefit of the use of mixed amines, but not to the point of making a revolutionary improvement in the cost of carbon capture.

Appendix A:

Figure A.1. Process Flow Diagram



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Appendix B: Stream Tables

Table B.1a. Stream Table: Baseline Scenario – 30% MEA Solvent.

Stream Name	FLUEGAS	COMPFLU	WATEROUT	FEEDGAS	TOWASH	CLEANER
Temperature °C	54.4	55	44.4	44.4	55.7	64
Pressure, bar	1.011	1.037	1.037	1.037	1.016	1.014
<i>Component Mole Flow, kmol/hr</i>						
CO ₂	3,300	3,300	-	3,300	280	280
H ₂ O	5,300	5,300	2,500	2,900	5,000	8,100
N ₂	23,288	23,288	-	23,288	23,288	23,288
O ₂	2,300	2,300	-	2,300	2,300	2,300
HCL	-	-	-	-	-	-
H ₃ O ⁺	-	-	-	-	-	-
HCO ₃ ⁻	-	-	-	-	-	-
CL ⁻	-	-	-	-	-	-
CO ₃ ⁻⁻	-	-	-	-	-	-
MEA	-	-	-	-	-	-
MEA ⁺	-	-	-	-	-	-
MEACOO ⁻	-	-	-	-	-	-
<i>Component Mass Flow, kg/hr</i>						
CO ₂	144,000	144,000	4	144,000	12,000	12,000
H ₂ O	97,200	96,000	44,000	52,000	90,000	145,000
N ₂	651,600	652,000	-	652,000	652,000	652,000
O ₂	72,000	72,000	-	72,000	72,000	72,000
HCL	-	-	-	-	-	-
H ₃ O ⁺	-	-	-	-	-	-
HCO ₃ ⁻	-	-	-	-	-	-
CL ⁻	-	-	-	-	-	-
CO ₃ ⁻⁻	-	-	-	-	-	-
MEA	-	-	-	-	-	-
MEA ⁺	-	-	-	-	-	-
MEACOO ⁻	-	-	-	-	-	-
<i>Total Flow, kmol/hr</i>	34,000	34,000	2,500	32,000	31,000	34,000
<i>Total Flow, kg/hr</i>	965,000	921,000	44,000	918,000	827,000	882,000
<i>Total Flow, cum/hr</i>	914,400	896,000	43	806,000	828,000	936,000
Vapor Fraction	1	1	0	1	1	1
Liquid Fraction	0	0	1	0	0	0

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Table B.1b. Stream Table: Baseline Scenario – 30% MEA Solvent.

Stream Name	AMINEFD	REC	COMBFD	OUTWASH	RICHOUT
Temperature °C	44.4	44.4	44.4	64	52.5
Pressure, bar	1.013	1.048	1.016	1.014	1.016
<i>Component Mole Flow, kmol/hr</i>					
CO2	-	-	-	-	4
H2O	1,800	121,000	123,000	510	121,000
N2	-	-	-	-	-
O2	-	-	-	-	-
HCL	-	-	-	-	-
H3O+	-	-	-	-	-
HCO3-	-	43	43	-	500
CL-	-	97	97	-	97
CO3--	-	40	40	-	29
MEA	58	6,500	6,600	-	1,000
MEA+	-	4,800	4,800	-	7,800
MEACOO-	-	4,600	4,600	-	7,100
<i>Component Mass Flow, kg/hr</i>					
CO2	-	-	-	-	130
H2O	32,400	2,176,000	2,209,000	9,300	2,172,000
N2	-	-	-	-	11
O2	-	-	-	-	4
HCL	-	-	-	-	-
H3O+	-	-	-	-	-
HCO3-	-	2,600	2,600	-	31,000
CL-	-	3,400	3,400	-	3,400
CO3--	-	2,400	2,400	-	1,700
MEA	3,500	397,000	400,000	-	63,000
MEA+	-	300,000	300,000	-	484,000
MEACOO-	-	480,000	480,000	-	743,000
<i>Total Flow, kmol/hr</i>	1,900	137,000	139,000	510	137,000
<i>Total Flow, kg/hr</i>	36,000	3,362,000	3,394,000	9,300	3,497,000
<i>Total Flow, cum/hr</i>	65	3,500	3,500	11	3,700
Vapor Fraction	0	0	0	0	0
Liquid Fraction	1	1	1	1	1

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Table B.1c. Stream Table: Baseline Scenario – 30% MEA Solvent.

Stream Name	STRIPFEE	LEANOUT	VAPOR	REFLUX	GASFSTR	CO2LIQ
Temperature °C	97.7	111.6	96.6	47.7	47.7	44.4
Pressure, bar	1.662	1.52	1.52	1.52	1.52	153
<i>Component Mole Flow, kmol/hr</i>						
CO2	970	11	3,000	-	3,000	3,000
H2O	121,000	121,000	2,800	2,500	240	11
N2	-	-	-	-	-	-
O2	-	-	-	-	-	-
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	510	240	-	4	-	-
CL-	97	97	-	-	-	-
CO3--	11	7	-	-	-	-
MEA	3,000	6,700	4	-	-	-
MEA+	6,800	4,800	-	4	-	-
MEACOO-	6,200	4,400	-	-	-	-
<i>Component Mass Flow, kg/hr</i>						
CO2	43,000	530	131,000	58	131,000	131,000
H2O	2,172,000	2,173,000	50,000	46,000	4,300	180
N2	11	-	11	-	11	11
O2	4	-	4	-	4	4
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	31,000	15,000	-	180	-	-
CL-	3,400	3,400	-	-	-	-
CO3--	620	480	-	-	-	-
MEA	181,000	410,000	180	-	-	-
MEA+	423,000	298,000	-	180	-	-
MEACOO-	643,000	463,000	-	-	-	-
<i>Total Flow, kmol/hr</i>	138,000	137,000	5,700	2,500	3,200	3,000
<i>Total Flow, kg/hr</i>	3,497,000	3,362,000	181,000	46,000	135,000	131,000
<i>Total Flow, cum/hr</i>	35,000	3,700	115,000	47	56,000	194
Vapor Fraction	0.012	0	1	0	1	1
Liquid Fraction	0.988	1	0	1	0	0

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Table B.1d. Stream Table: Baseline Scenario – 30% MEA Solvent.

Stream Name	LEANTOHX	LEANFRHX	RICHTOHX
Temperature °C	111.7	57	52.6
Pressure, bar	1.703	1.703	4.42
<i>Component Mole Flow, kmol/hr</i>			
CO2	11	-	4
H2O	121,000	121,000	121,000
N2	-	-	-
O2	-	-	-
HCL	-	-	-
H3O+	-	-	-
HCO3-	238	65	500
CL-	97	97	97
CO3--	7	29	29
MEA	6,700	6,500	1,000
MEA+	4,800	4,800	7,800
MEACOO-	4,400	4,600	7,100
<i>Component Mass Flow, kg/hr</i>			
CO2	530	4	130
H2O	2,173,000	2,176,000	2,172,000
N2	-	-	11
O2	-	-	4
HCL	-	-	-
H3O+	-	-	-
HCO3-	15,000	3,900	31,000
CL-	3,400	3,400	3,400
CO3--	480	1,800	1,700
MEA	410,000	398,000	63,000
MEA+	298,000	300,000	484,000
MEACOO-	463,000	480,000	743,000
<i>Total Flow, kmol/hr</i>	137,000	137,000	137,000
<i>Total Flow, kg/hr</i>	3,362,000	3,362,000	3,497,000
<i>Total Flow, cum/hr</i>	3,700	3,500	3,700
Vapor Fraction	0	0	0
Liquid Fraction	1	1	1

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Table B.2a. Stream Table: 30% Mixed Amine Solvent.

Stream Name	FLUEGAS	COMPFLU	WATEROUT	FEEDGAS	TOWASH	CLEANER
Temperature °C	54.4	55	44.4	44.4	50.6	50.3
Pressure, bar	1.011	1.037	1.037	1.037	1.016	1.014
<i>Component Mole Flow, kmol/hr</i>						
CO2	3,300	3,300	-	3,300	300	300
H2O	5,300	5,300	2,462	2,900	3,700	3,700
N2	23,000	23,000	-	23,000	23,000	23,000
O2	2,300	2,300	-	2,300	2,300	2,300
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	-	-	-	-	-	-
CL-	-	-	-	-	-	-
CO3--	-	-	-	-	-	-
MEA	-	-	-	-	-	-
MEA+	-	-	-	-	-	-
MEACOO-	-	-	-	-	-	-
MDEA	-	-	-	-	-	-
AMP	-	-	-	-	-	-
AMP+	-	-	-	-	-	-
MDEA+	-	-	-	-	-	-
<i>Component Mass Flow, kg/hr</i>						
CO2	144,000	144,000	-	144,000	13,000	13,000
H2O	96,000	96,000	44,000	52,000	67,000	66,000
N2	652,000	652,000	-	652,000	652,000	652,000
O2	72,000	72,000	-	72,000	72,000	72,000
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	-	-	-	-	-	-
CL-	-	-	-	-	-	-
CO3--	-	-	-	-	-	-
MEA	-	-	-	-	-	-
MEA+	-	-	-	-	-	-
MEACOO-	-	-	-	-	-	-
MDEA	-	-	-	-	-	-
AMP	-	-	-	-	-	-
AMP+	-	-	-	-	-	-
MDEA+	-	-	-	-	-	-
<i>Total Flow, kmol/hr</i>	34,000	34,000	2,500	32,000	30,000	30,000
<i>Total Flow, kg/hr</i>	965,000	965,000	44,000	921,000	805,000	804,000
<i>Total Flow, cum/hr</i>	914,000	896,000	43	806,000	785,000	781,000
Vapor Fraction	1	1	0	1	1	1
Liquid Fraction	0	0	1	0	0	0

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Table B.2b. Stream Table: 30% Mixed Amine Solvent.

Stream Name	AMINEFD	REC	COMBFD	OUTWASH	RICHOUT
Temperature °C	44.4	44.4	44.4	50.3	50
Pressure, bar	1.013	1.048	1.016	1.014	1.016
<i>Component Mole Flow, kmol/hr</i>					
CO2	-	-	-	-	4
H2O	4,600	140,000	137,000	3,700	139,000
N2	-	-	-	-	-
O2	-	-	-	-	-
HCL	-	-	-	-	-
H3O+	-	-	-	-	-
HCO3-	-	68	68	-	1,224
CL-	-	-	-	-	-
CO3--	-	29	29	-	72
MEA	-	2,400	2,400	-	170
MEA+	-	2,900	2,900	-	3,300
MEACOO-	-	3,700	3,700	-	5,400
MDEA	-	3,300	3,300	-	1,300
AMP	-	620	620	-	86
AMP+	-	600	600	-	1,100
MDEA+	-	350	350	-	2,400
<i>Component Mass Flow, kg/hr</i>					
CO2	-	-	-	-	160
H2O	84,000	2,516,000	2,470,000	66,000	2,499,000
N2	-	-	-	-	14
O2	-	-	-	-	4
HCL	-	-	-	-	-
H3O+	-	-	-	-	-
HCO3-	-	4,300	4,100	-	74,700
CL-	-	-	-	-	-
CO3--	-	1,800	1,700	-	4,400
MEA	-	145,000	145,000	-	11,000
MEA+	-	177,000	177,000	-	204,000
MEACOO-	-	382,000	382,000	-	565,000
MDEA	-	392,000	392,000	-	152,000
AMP	-	55,000	55,000	-	7,600
AMP+	-	54,000	54,000	-	102,000
MDEA+	-	42,000	42,000	-	285,000
<i>Total Flow, kmol/hr</i>	4,600	154,000	151,000	3,700	154,000
<i>Total Flow, kg/hr</i>	84,000	3,771,000	3,724,000	66,000	3,905,000
<i>Total Flow, cum/hr</i>	65	3,800	3,800	68	3,900
Vapor Fraction	0	0	0	0	0
Liquid Fraction	1	1	1	1	1

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Table B.2c. Stream Table: 30% Mixed Amine Solvent.

Stream Name	STRIPFEE	LEANOUT	VAPOR	REFLUX	GASFSTR	CO2LIQ
Temperature °C	99.4	110.6	98	47.7	47.7	44.4
Pressure, bar	1.662	1.52	1.52	1.52	1.52	153
<i>Component Mole Flow, kmol/hr</i>						
CO2	960	7	2,952	-	2,900	2,900
H2O	139,000	140,000	3,700	3,500	230	230
N2	-	-	-	-	-	-
O2	-	-	-	-	-	-
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	688	227	-	-	-	-
CL-	-	-	-	-	-	-
CO3--	14	7	-	-	-	-
MEA	810	2,900	-	-	-	-
MEA+	3,000	2,500	-	-	-	-
MEACOO-	5,100	3,500	-	-	-	-
MDEA	1,800	2,800	-	-	-	-
AMP	300	730	-	-	-	-
AMP+	920	490	-	-	-	-
MDEA+	1,900	806	-	-	-	-
<i>Component Mass Flow, kg/hr</i>						
CO2	42,000	360	130,000	79	130,000	130,000
H2O	2,510,000	2,514,000	67,000	62,000	4,200	180
N2	14	-	14	-	14	14
O2	4	-	4	-	4	4
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	41,969	13,939	-	79	-	-
CL-	-	-	-	-	-	-
CO3--	950	380	-	-	-	-
MEA	50,000	176,000	54	-	-	-
MEA+	187,000	154,000	-	54	-	-
MEACOO-	527,000	368,000	-	-	-	-
MDEA	214,000	339,000	36	-	-	-
AMP	27,000	65,000	11	-	-	-
AMP+	83,000	44,000	-	11	-	-
MDEA+	223,000	97,000	-	36	-	-
<i>Total Flow, kmol/hr</i>	138,000	154,000	7,000	3,500	3,200	3,000
<i>Total Flow, kg/hr</i>	3,497,000	3,771,000	197,000	63,000	134,000	130,000
<i>Total Flow, cum/hr</i>	35,000	4,000	134,000	65	56,000	194
Vapor Fraction	0.012	0	1	0	1	1
Liquid Fraction	0.988	1	0	1	0	0

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Table B.2d. Stream Table: 30% Mixed Amine Solvent.

Stream Name	LEANTOHX	LEANFRHX	RICHTOHX
Temperature °C	110.6	54.4	50
Pressure, bar	1.703	1.703	442
<i>Component Mole Flow, kmol/hr</i>			
CO2	7	-	4
H2O	140,000	140,000	139,000
N2	-	-	-
O2	-	-	-
HCL	-	-	-
H3O+	-	-	-
HCO3-	227	86	1,224
CL-	-	-	-
CO3--	7	25	72
MEA	2,900	2,400	170
MEA+	2,500	2,800	3,300
MEACOO-	3,500	3,700	5,400
MDEA	2,800	3,200	1,300
AMP	730	640	86
AMP+	490	580	1,100
MDEA+	810	420	2,400
<i>Component Mass Flow, kg/hr</i>			
CO2	360	4	160
H2O	2,514,000	2,516,000	2,499,000
N2	-	-	14
O2	-	-	4
HCL	-	-	-
H3O+	-	-	-
HCO3-	14,000	5,300	75,000
CL-	-	-	-
CO3--	380	1,400	4,400
MEA	176,000	149,000	11,000
MEA+	154,000	174,000	204,000
MEACOO-	368,000	382,000	565,000
MDEA	339,000	384,000	152,000
AMP	65,000	57,000	7,600
AMP+	44,000	53,000	102,000
MDEA+	97,000	51,000	285,000
<i>Total Flow, kmol/hr</i>	154,000	154,000	154,000
<i>Total Flow, kg/hr</i>	3,771,000	3,771,000	3,905,000
<i>Total Flow, cum/hr</i>	4,000	3,900	3,900
Vapor Fraction	0	0	0
Liquid Fraction	1	1	1

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Table B.3a. Stream Table: 35% Mixed Amine Solvent.

Stream Name	FLUEGAS	COMPFLU	WATEROUT	FEEDGAS	TOWASH	CLEANER
Temperature °C	54.4	55	44.4	44.4	51.5	51.2
Pressure, bar	1.011	1.037	1.037	1.037	1.016	1.014
<i>Component Mole Flow, kmol/hr</i>						
CO2	3,300	3,300	-	3,300	300	300
H2O	5,300	5,300	2,500	2,900	3,900	3,900
N2	23,000	23,000	-	23,000	23,000	23,000
O2	2,300	2,300	-	2,300	2,300	2,300
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	-	-	-	-	-	-
CL-	-	-	-	-	-	-
CO3--	-	-	-	-	-	-
MEA	-	-	-	-	-	-
MEA+	-	-	-	-	-	-
MEACOO-	-	-	-	-	-	-
MDEA	-	-	-	-	-	-
AMP	-	-	-	-	-	-
AMP+	-	-	-	-	-	-
MDEA+	-	-	-	-	-	-
<i>Component Mass Flow, kg/hr</i>						
CO2	144,000	144,000	-	144,000	14,000	14,000
H2O	96,000	96,000	44,000	52,000	71,000	70,000
N2	652,000	652,000	-	652,000	652,000	652,000
O2	72,000	72,000	-	72,000	72,000	72,000
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	-	-	-	-	-	-
CL-	-	-	-	-	-	-
CO3--	-	-	-	-	-	-
MEA	-	-	-	-	-	-
MEA+	-	-	-	-	-	-
MEACOO-	-	-	-	-	-	-
MDEA	-	-	-	-	-	-
AMP	-	-	-	-	-	-
AMP+	-	-	-	-	-	-
MDEA+	-	-	-	-	-	-
<i>Total Flow, kmol/hr</i>	34,000	34,000	2,500	32,000	30,000	30,000
<i>Total Flow, kg/hr</i>	965,000	965,000	44,000	921,000	810,000	809,000
<i>Total Flow, cum/hr</i>	914,000	896,000	43	806,000	792,000	790
Vapor Fraction	1	1	0	1	1	1
Liquid Fraction	0	0	1	0	0	0

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Table B.3b. Stream Table: 35% Mixed Amine Solvent.

Stream Name	AMINEFD	REC	COMBFD	OUTWASH	RICHOUT
Temperature °C	44.4	44.4	44.4	51.2	49.9
Pressure, bar	1.013	1.048	1.016	1.014	1.016
<i>Component Mole Flow, kmol/hr</i>					
CO2	-	-	-	-	4
H2O	4,800	111,000	108,000	3,700	110,000
N2	-	-	-	-	-
O2	-	-	-	-	-
HCL	-	-	-	-	-
H3O+	-	-	-	-	-
HCO3-	-	50	50	-	960
CL-	-	-	-	-	-
CO3--	-	22	18	-	54
MEA	-	2,500	2,500	-	170
MEA+	-	2,800	2,800	-	3,100
MEACOO-	-	3,600	3,600	-	5,600
MDEA	-	3,300	3,300	-	1,200
AMP	-	630	630	-	80
AMP+	-	590	590	-	1,100
MDEA+	-	350	350	-	2,400
<i>Component Mass Flow, kg/hr</i>					
CO2	-	-	-	-	120
H2O	87,000	1,995,000	1,953,000	66,000	1,983,000
N2	-	-	-	-	11
O2	-	-	-	-	4
HCL	-	-	-	-	-
H3O+	-	-	-	-	-
HCO3-	-	3,200	3,100	-	59,000
CL-	-	-	-	-	-
CO3--	-	1,200	1,200	-	3,200
MEA	-	155,000	155,000	-	11,000
MEA+	-	171,000	171,000	-	194,000
MEACOO-	-	375,000	375,000	-	583,000
MDEA	-	393,000	393,000	-	147,000
AMP	-	56,000	56,000	-	7,200
AMP+	-	53,000	53,000	-	103,000
MDEA+	-	42,000	42,000	-	291,000
<i>Total Flow, kmol/hr</i>	4,800	125,000	122,000	3,700	125,000
<i>Total Flow, kg/hr</i>	87,000	3,245,000	3,202,000	66,000	3,379,000
<i>Total Flow, cum/hr</i>	65	3,300	3,300	68	3,400
Vapor Fraction	0	0	0	0	0
Liquid Fraction	1	1	1	1	1

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Table B.3c. Stream Table: 35% Mixed Amine Solvent.

Stream Name	STRIPFEE	LEANOUT	VAPOR	REFLUX	GASFSTR	CO2LIQ
Temperature °C	99.6	111.3	98.2	47.7	47.7	44.4
Pressure, bar	1.662	1.52	1.52	1.52	1.52	153
<i>Component Mole Flow, kmol/hr</i>						
CO2	910	7	3,000	-	2,900	2,900
H2O	111,000	111,000	3,500	3,200	230	11
N2	-	-	-	-	-	-
O2	-	-	-	-	-	-
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	500	170	-	-	-	-
CL-	-	-	-	-	-	-
CO3--	11	4	-	-	-	-
MEA	830	3,000	-	-	-	-
MEA+	2,900	2,400	-	-	-	-
MEACOO-	5,200	3,500	-	-	-	-
MDEA	1,700	2,800	-	-	-	-
AMP	290	730	-	-	-	-
AMP+	930	490	-	-	-	-
MDEA+	1,900	817	-	-	-	-
<i>Component Mass Flow, kg/hr</i>						
CO2	40,000	330	130,000	72	130,000	130,000
H2O	1,992,000	1,994,000	62,000	58,000	4,200	176
N2	11	-	11	-	11	11
O2	4	-	4	-	4	4
HCL	-	-	-	-	-	-
H3O+	-	-	-	-	-	-
HCO3-	31,000	10,000	-	108	-	-
CL-	-	-	-	-	-	-
CO3--	670	260	-	-	-	-
MEA	51,000	185,000	76	-	-	-
MEA+	178,000	147,000	-	76	-	-
MEACOO-	541,000	364,000	-	-	-	-
MDEA	205,000	337,000	47	-	-	-
AMP	26,000	65,000	11	-	-	-
AMP+	84,000	44,000	-	11	-	-
MDEA+	231,000	98,000	-	47	-	-
<i>Total Flow, kmol/hr</i>	126,000	125,000	6,400	3,200	3,200	3,000
<i>Total Flow, kg/hr</i>	3,379,000	3,245,000	193,000	59,000	134,000	130,000
<i>Total Flow, cum/hr</i>	37,000	3,400	130,000	58	56,000	190
Vapor Fraction	0.015	0	1	0	1	1
Liquid Fraction	0.985	1	0	1	0	0

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Table B.3d. Stream Table: 35% Mixed Amine Solvent.

Stream Name	LEANTOHX	LEANFRHX	RICHTOHX
Temperature °C	111.3	54.4	50
Pressure, bar	1.703	1.703	442
<i>Component Mole Flow, kmol/hr</i>			
CO2	7	-	4
H2O	111,000	111,000	110,000
N2	-	-	-
O2	-	-	-
HCL	-	-	-
H3O+	-	-	-
HCO3-	170	60	960
CL-	-	-	-
CO3--	4	18	54
MEA	3,000	2,600	200
MEA+	2,400	2,700	3,100
MEACOO-	3,500	3,600	5,600
MDEA	2,800	3,200	1,200
AMP	730	640	79
AMP+	490	580	1,100
MDEA+	820	420	2,400
<i>Component Mass Flow, kg/hr</i>			
CO2	330	4	120
H2O	1,994,000	1,995,000	1,983,000
N2	-	-	11
O2	-	-	4
HCL	-	-	-
H3O+	-	-	-
HCO3-	10,000	4,000	59,000
CL-	-	-	-
CO3--	260	980	3,200
MEA	185,000	159,000	11,000
MEA+	147,000	167,000	194,000
MEACOO-	364,000	374,000	583,000
MDEA	337,000	385,000	146,000
AMP	65,000	57,000	7,200
AMP+	44,000	52,000	103,000
MDEA+	98,000	50,000	291,000
<i>Total Flow, kmol/hr</i>	125,000	123,000	125,000
<i>Total Flow, kg/hr</i>	3,245,000	3,245,000	3,379,000
<i>Total Flow, cum/hr</i>	3,400	3,300	3,400
Vapor Fraction	0	0	0
Liquid Fraction	1	1	1

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Appendix C: Equipment Lists for 30% Mixed Amine and 35% Mixed Amines

Table C.1. 30% Mixed Amine Major Process Equipment List.

Item Label in Process Flow Diagram (PFD)	Number	Size	Description	Comments
Columns				
ABSORBER	2	25 m ht x 8.6 m dia	304 SS column shell 15 m of 316 SS structured packing	Vapor wash step at top of column to reduce amine emissions is modeled as COND
STRIPPER	2	10 m ht x 4.8 m dia	304 SS column 6 m of 316 SS Structured packing	
Pressure Vessels				
DRYER	2	7 m x 3.5 m dia	Knock-out drum for stripper	304 SS
COND1	1	5 m x 2.5 m dia	First CO ₂ compression stage knock-out drum	304 SS
COND2	1	3.0 x 1.5 m dia	Second CO ₂ compression stage knock-out drum	304 SS
COND3	1	1.8 x 0.9 m dia	Third CO ₂ compression stage knock-out drum	304 SS
COND4	1	1.1 x 0.6 m dia	Fourth CO ₂ compression stage knock-out drum	CS
COND5	1	0.6 x 0.3 m dia	Fifth CO ₂ compression stage knock-out drum	CS
Heat Exchangers				
REGEN	14	1,000 m ²	770 GJ/hr exchange	Shell: 304 SS Tubes: 304 SS
DRYER	2	1,000 m ²	-160 GJ/hr cooling	Shell: CS Tubes: 304 SS
REBOIL	4	1,000 m ²	415 GJ/hr heating	Shell: CS Tubes: 304 SS
COOLER	4	1,000 m ²	-137 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND1	4	900 m ²	-17 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND2	3	800 m ²	-11 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND3	3	800 m ²	-11 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND4	3	800 m ²	-13 GJ/hr cooling	Shell: CS Tubes: CS
CO2COND5	5	1,000 m ²	-24 GJ/hr cooling	Shell: CS Tubes: 304 SS
PRECOOLERHX	10	1,000 m ²	-113 GJ/hr	Shell: CS Tubes: 304 SS Not shown on flow diagram
INTERCOOLER	2	1,000 m ²	-72 GJ/hr	Shell: CS Tubes: 304 SS Not shown on flow diagram

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Item Label in Process Flow Diagram (PFD)	Number	Size	Description	Comments
Compressors				
COMP	7	0.1 MW	Pre-Absorber Blower	Stainless Steel
COMP1	1	2.25 MW	Brake Power	Stainless Steel
COMP2	1	2.25 MW	Brake Power	Stainless Steel
COMP3	1	2.25 MW	Brake Power	Stainless Steel
COMP4	1	2.25 MW	Brake Power	CS
COMP5	1	2.25 MW	Brake Power	CS
Pumps				
Absorber Bottoms Pump	5	0.275 m ³ /s	750 kPa ΔP	304 SS
Stripper Bottoms Pump	5	0.275 m ³ /s	300 kPa ΔP	304 SS
Intercooling Pump	4	0.275 m ³ /s	30 kPa ΔP	304 SS. Not shown on flow diagram
Tankage				
Amine Solution Surge Tank	2	175 m ³		304 SS
Amine Solution Mixing Tank	1	90 m ³		304 SS
Caustic Mixing Tank	1	10 m ³		Glass-lined CS.
Caustic Storage Tank	1	400 m ³		Glass-lined CS. 28 day storage
MEA Feed Storage Tank	1	80 m ³		304 SS. 28 day storage
MDEA Feed Storage Tank	1	20 m ³		304 SS. 28 day storage
AMP Feed Storage Tank	1	3 m ³		304 SS. 28 day storage
Other				
Ion Exchange Column	2	7.75 m x 2.5 m dia	Used to remove heat-stable salts from the amine solution	Resin-lined CS pressure vessel. 1 hour space velocity. Not shown on flow diagram
Activated Carbon Beds for Amine Solution	2	6 m x 2 m dia	Used to remove amine degradation byproducts from amine solution	Activated carbon bed with resin-lined CS. 0.5 hour space velocity. Not shown on flow diagram
CO2 Drying Beds	2	5.5 m x 1.5 m dia	Used to dry CO2 after COND3 knock-out drum	Silica: CS vessel shell. Not shown on flow diagram
Activated Carbon Beds for treatment of CO2	2	4 m x 1 m dia	For removal of organic impurities from CO ₂ .	Activated carbon with resin-lined CS. Not shown on flow diagram

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Table C.2. 35% Mixed Amine Major Process Equipment List.

Item Label in Process Flow Diagram (PFD)	Number	Size	Description	Comments
Columns				
ABSORBER	2	25 m ht x 8.5 m dia	304 SS column shell 15 m of 316 SS structured packing	Vapor wash step at top of column to reduce amine emissions is modeled as COND
STRIPPER	2	10 m ht x 4.8 m dia	304 SS column 6 m of 316 SS Structured packing	
Pressure Vessels				
DRYER	2	6.75 m x 3.5 m dia	Knock-out drum for stripper	304 SS
COND1	1	5 m x 2.5 m dia	First CO ₂ compression stage knock-out drum	304 SS
COND2	1	3.0 x 1.5 m dia	Second CO ₂ compression stage knock-out drum	304 SS
COND3	1	1.8 x 0.9 m dia	Third CO ₂ compression stage knock-out drum	304 SS
COND4	1	1.1 x 0.6 m dia	Fourth CO ₂ compression stage knock-out drum	CS
COND5	1	0.6 x 0.3 m dia	Fifth CO ₂ compression stage knock-out drum	CS
Heat Exchangers				
REGEN	13	1,000 m ²	650 GJ/hr exchange	Shell: 304 SS Tubes: 304 SS
DRYER	2	900 m ²	-150 GJ/hr cooling	Shell: CS Tubes: 304 SS
REBOIL	4	1,000 m ²	392 GJ/hr heating	Shell: CS Tubes: 304 SS
COOLER	4	1,000 m ²	-115 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND1	4	900 m ²	-17 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND2	3	800 m ²	-11 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND3	3	800 m ²	-11 GJ/hr cooling	Shell: CS Tubes: 304 SS
CO2COND4	3	800 m ²	-13 GJ/hr cooling	Shell: CS Tubes: CS
CO2COND5	5	1,000 m ²	-24 GJ/hr cooling	Shell: CS Tubes: 304 SS
PRECOOLERHX	10	1,000 m ²	-113 GJ/hr	Shell: CS Tubes: 304 SS Not shown on flow diagram
INTERCOOLER	2	1,000 m ²	-72 GJ/hr	Shell: CS Tubes: 304 SS Not shown on flow diagram

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Item Label in Process Flow Diagram (PFD)	Number	Size	Description	Comments
Compressors				
COMP	7	0.1 MW	Pre-Absorber Blower	Stainless Steel
COMP1	1	2.25 MW	Brake Power	Stainless Steel
COMP2	1	2.25 MW	Brake Power	Stainless Steel
COMP3	1	2.25 MW	Brake Power	Stainless Steel
COMP4	1	2.25 MW	Brake Power	CS
COMP5	1	2.25 MW	Brake Power	CS
Pumps				
Absorber Bottoms Pump	5	0.245 m ³ /s	750 kPa ΔP	304 SS
Stripper Bottoms Pump	5	0.245 m ³ /s	300 kPa ΔP	304 SS
Intercooling Pump	4	0.245 m ³ /s	30 kPa ΔP	304 SS. Not shown on flow diagram
Tankage				
Amine Solution Surge Tank	2	140 m ³		304 SS
Amine Solution Mixing Tank	1	65 m ³		304 SS
Caustic Mixing Tank	1	10 m ³		Glass-lined CS.
Caustic Storage Tank	1	400 m ³		Glass-lined CS. 28 day storage
MEA Feed Storage Tank	1	80 m ³		304 SS. 28 day storage
MDEA Feed Storage Tank		20 m ³		304 SS. 28 day storage
AMP Feed Storage Tank		3 m ³		304 SS. 28 day storage
Other				
Ion Exchange Column	2	7.2 m x 2.4 m dia	Used to remove heat-stable salts from the amine solution	Resin-lined CS pressure vessel. 1 hour space velocity. Not shown on flow diagram
Activated Carbon Beds for Amine Solution	2	5.75 m x 2 m dia	Used to remove amine degradation byproducts from amine solution	Activated carbon bed with resin-lined CS. 0.5 hour space velocity. Not shown on flow diagram
CO ₂ Drying Beds	2	5.5 m x 1.5 m dia	Used to dry CO ₂ after COND3 knock-out drum	Silica: CS vessel shell. Not shown on flow diagram
Activated Carbon Beds for treatment of CO ₂	2	4 m x 1 m dia	For removal of organic impurities from CO ₂ .	Activated carbon with resin-lined CS. Not shown on flow diagram

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Appendix D: Detailed Cost Estimate Tables

Table D.1. Capital Cost Estimate: Baseline Scenario – 30% MEA Solvent.

Table D.1
CO₂ Removal Using 30% MEA, Gibson 3,
Annual Estimated Variable Costs

Costs in 2011 US\$

Onstream Factor 0.8

Losses to Energy Generation

Lost Generation From Steam Bleed to Reboiler	28.1 MWh	\$51.2 per MWh*	\$1,439	\$10,083,000
Power generated by Letdown Turbine	(5.4) MWh	\$51.2 per MWh*	\$(274)	\$(1,923,000)
Parasitic Electricity Consumption by Compressors/Pumps	16.6 MWh	\$51.2 per MWh*	\$ 847	\$ 5,938,000
Losses to Energy Generation			\$ 2,012	\$ 14,098,000

Materials Consumed	Number/hr	Unit Cost \$	Costs/hr	Costs/yr
Monoethanolamine	0.21 tonnes	\$2,880 per tonne	\$ 603	\$ 4,227,000
Stabilizer (5% of Amine makeup costs)				\$211,000
Caustic Soda	0.16 tonnes	\$460 per tonne	\$74	\$516,000
Desiccant Replacement	0.0031 tonnes	\$5,000 per tonne	\$15	\$107,000
Activated Carbon	0.0024 tonnes	\$2,500 per tonne	\$6	\$43,000
Ion Exchange Resin	0.0059 tonnes	\$8,000 per tonne	\$47	\$333,000
Disposal/treatment of Selenium-containing wastes	0.1140 tonnes	\$300 per tonne	\$34	\$240,000
Other Waste Disposal	0.0650 tonnes	\$300 per tonne	\$20	\$137,000
Material Consumed Costs			\$799	\$5,814,000
Utilities				
Cooling Water	25,920 cu.m	\$0.02 per cu.m	\$518	\$3,633,000
Utility Costs			\$518	\$3,633,000
Total Variable costs			\$3,329	\$23,545,000

* 2006-2011 Five-year average bilateral contract price for Cinergy/MISO hub used as the cost of electricity

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Table D.2. Variable Cost Estimate: Baseline Scenario – 30% MEA Solvent.

Table D.2 CO ₂ Removal Using 30% MEA, Gibson 3, Estimated Fixed Capital Costs			
Plant Net Capacity, CO2 Removed/Hr		131 tonnes/hr 144 short tons/hr	
Costs in 2011 thousand US\$			
Gas Cleanup and Power Plant			
Battery Limits Investment	Equipment Cost	Installation Cost	Total Cost
Absorber Columns	\$4,380	\$9,640	\$14,020
Stripper Columns	\$1,080	\$2,370	\$3,450
Heat Exchangers	\$7,600	\$11,900	\$19,500
Pressure Vessels	\$2,450	\$3,430	\$5,880
Fans	\$ 175	\$245	\$420
CO2 Compressors	\$6,260	\$9,350	\$15,610
Pumps	\$1,660	\$3,920	\$5,580
Let-down Turbine	\$1,800	\$1,200	\$3,000
Subtotal			\$67,460
BLI Contingency	20%	of Installed Equipment Costs	\$13,490
Battery Limits Investment			\$80,950
Tankage	Equipment Cost	Installation Cost	
Amine Solution Surge Tanks	\$260	\$310	\$570
Feed Amine Storage Tank	\$110	\$130	\$240
Makeup Amine Mix/Staging Tank	\$100	\$120	\$220
Caustic Soda Tank	\$120	\$210	\$330
Tankage Investment Subtotal			\$1,360
Utilities	Equipment Cost	Installation Cost	Investment
Cooling Water	\$3,840	\$ 2,610	\$6,450
Utilities Investment Subtotal			\$6,500
Pipelines from Main Plant	Unit cost, installed	Length (km)	Diameter (cm)
Flue Gas	\$16	0.5	660
Steam	\$16	0.5	60
Condensate	\$16	0.5	80
Pipeline Investment Subtotal			\$6,400
Offsites & Utility Investment Contingency		20%	\$2,852
Offsite & Utilities Investment			\$17,112
General Service Facilities	10%	of BLI & Utilities Investment	\$9,810
Waste Treatment	5%	of BLI Investment	\$4,050
Outside Battery Limits Investment			\$31,000
Greenfield Total Plant Investment			\$112,000
Retrofit Complexity Factor	20%	Additional Retrofit Costs	\$22,400
Total Estimated Investment			\$134,400

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Table D.3. Operating Cost Estimate: Baseline Scenario – 30% MEA Solvent.

Table D.3 CO2 Removal Using 30% MEA, Gibson 3, Annual Estimated Operating Costs

Plant Net Capacity	0.92 megatonnes/yr 1.01 MMshort tons/yr
Onstream factor	0.8
Price of Electricity	\$51.20 per MWh

Costs in 2011 thousand US\$ unless noted

	Costs
Plant Investment, Battery Limits (BLI)	\$97,100
Plant Investment, Outside Battery Limits (OBLI)	\$37,200
Total Fixed Capital (TFM)	\$134,300
Operating Costs, Per Year	
Variable Costs	
Losses to Energy Generation (net)	\$14,098
Materials Consumed Costs	\$5,810
Utility Costs (net)	\$3,630
Total Annual	\$23,538
Labor Costs	
Operating Labor (3 shifts)	6 \$62 wages/year \$372
Maintenance Labor	3.00% of BLI \$2,913
Control Laboratory Labor, 10% of Operating Labor	10% of Operating Labor \$40
Direct Labor Costs	\$3,330
Maintenance Materials	3.00% of BLI \$2,910
Operating Supplies, 12% of Operating Labor	12% of Operating Labor \$40
Total Direct Costs	\$29,818
Plant Overhead	80% of Direct Labor Costs \$2,660
Taxes and Insurance	1.60% of TFC \$2,150
Cash Costs	\$34,628
Levelized Depreciation	3.3% of TFC \$4,480
Gate Costs	\$39,108
General, Admin, Sales, Research	5% of Gate Costs \$1,960
Production Costs, excluding Cost of Capital	\$41,100
TFC + Estimated Working Capital*	\$143,500
Annual Cost of Capital	12.5% of Capital Investment \$17,940
Total Operating Costs	\$59,040

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<i>Cost per ton CO2, dollars</i>	<i>\$58.40</i>
<i>Cost per tonne CO2, dollars</i>	<i>\$64.40</i>
Additional costs per MWh	\$68.7 MWh
Nominal Output of Gibson-3, MW	696 MWe
Nominal Output after existing peripheral losses, MW	635 MWe
Thermal Rating of Gibson-3, MW	1809 MWth
Share of Gibson-3 CO2 emissions processed	25.5% MW
Equivalent electricity output of share of plant emissions processed before carbon capture losses	162 MWe
Power losses from Carbon Capture and Compression, MW	39 MWe
Power output of share of plant for which emissions are processed after energy losses	123 MWe
Reduction in Power Output due to carbon capture	24.3%
Efficiency without Carbon Capture	35.1%
Efficiency with Carbon Capture	26.6%
Reduction in Efficiency	-8.50%
Annual Electrical Energy output of share of plant for which emissions are processed after carbon capture losses	859,952 MWh

* Working Capital was estimated at 25% of Annual Cash Costs plus cost of chemical and resin inventory

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Table D.4. Capital Cost Estimate: 30% Mixed Amine Solvent.

Table D.4
CO2 Removal Using 30% Mixed Amine (0.5 MEA, 0.4 MDEA, 0.1 AMP by weight), Gibson 3,
Annual Estimated Variable Costs

Costs in 2011 US\$

Onstream Factor 0.8

Losses to Energy Generation

Lost Generation From Steam Bleed to Reboiler	26.2 MWh	\$51 per MWh*	\$1,339	\$9,386,000
Power generated by Letdown Turbine	(4.7) MWh	\$51 per MWh*	\$(242)	\$(1,697,000)
Parasitic Electricity Consumption by Compressors/Pumps	16.6 MWh	\$51 per MWh*	\$847	\$ 5,938,000
Losses to Energy Generation			\$ 1,945	\$ 13,627,000

Materials Consumed	Number/hr	Unit Cost \$	Costs/hr	Costs/yr
Monoethanolamine (MEA)	0.11 tonnes	\$2,880 per tonne	\$302	\$2,119,000
Methyl Diethanolamine (MDEA)	0.024 tonnes	\$2,980 per tonne	\$70	\$491,000
2-Amino Methyl Propanol (AMP)	0.005 tonnes	\$4,960 per tonne	\$23	\$164,000
2-Amino Methyl Propanol (AMP)				\$139,000
Caustic Soda	0.16 tonnes	\$460 per tonne	\$74	\$516,000
Desiccant Replacement	0.0031 tonnes	\$5,000 per tonne	\$15	\$107,000
Activated Carbon	0.0024 tonnes	\$2,500 per tonne	\$6	\$43,000
Ion Exchange Resin	0.0059 tonnes	\$8,000 per tonne	\$47	\$333,000
Disposal/treatment of Selenium-containing wastes	0.1140 tonnes	\$300 per tonne	\$34	\$240,000
Other Waste Disposal	0.0650 tonnes	\$300 per tonne	\$20	\$137,000
Material Consumed Costs			\$593	\$4,289,000
Utilities				
Cooling Water	25,920 cu.m	\$0.02 per cu.m	<u>\$529</u>	<u>\$3,709,000</u>
Utility Costs			\$529	\$3,709,000
Total Variable costs			\$3,066	\$21,625,000

* 2006-2011 Five-year average bilateral contract price for Cinergy/MISO hub used as the cost of electricity

* Relative degradation rates of the individual amines were based on Ind. Eng. Chem. Res., 2009, 48 (20), pp 9061–9067: DOI: 10.1021/ie900472x

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Table D.5. Variable Cost Estimate: 30% Mixed Amine Solvent.

Table D.5 CO2 Removal Using 30% Mixed Amine (0.5 MEA, 0.4 MDEA, 0.1 AMP by weight), Gibson 3, Estimated Fixed Capital Costs			
Plant Net Capacity, CO2 Removed/Hr			130 tonnes/hr 144 short tons/hr
Costs in 2011 thousand US\$			
Gas Cleanup and Power Plant			
Battery Limits Investment	Equipment Cost	Installation Cost	Total Cost
Absorber Columns	\$4,220	\$9,290	\$13,510
Stripper Columns	\$1,080	\$2,380	\$3,460
Heat Exchangers	\$8,200	\$12,600	\$20,800
Pressure Vessels	\$2,550	\$3,570	\$6,120
Fans	\$ 175	\$245	\$420
CO2 Compressors	\$6,260	\$9,350	\$15,610
Pumps	\$1,660	\$3,920	\$5,580
Let-down Turbine	\$1,980	\$1,320	\$3,300
Subtotal			\$68,860
BLI Contingency	20%	of Installed Equipment Costs	\$13,760
Battery Limits Investment			\$82,560
Tankage	Equipment Cost	Installation Cost	
Amine Solution Surge Tanks	\$260	\$310	\$570
MEA Storage Tank	\$100	\$120	\$220
MDEA Storage Tank	\$60	\$70	\$130
AMP Storage Tank	\$40	\$50	\$90
Makeup Amine Staging Tank	\$100	\$120	\$220
Makeup Amine Staging Tank	\$100	\$120	\$220
Caustic Soda Tank	\$130	\$210	\$340
Tankage Investment Subtotal			\$1,790
Utilities	Equipment Cost	Installation Cost	Investment
Cooling Water	\$3,170	\$2,380	\$5,550
Utilities Investment Subtotal			\$5,600
Pipelines from Main Plant	Unit cost, installed	Length (km)	Diameter (cm)
Flue Gas	\$16/km-cm	0.5	660
Steam	\$16/km-cm	0.5	60
Condensate	\$16/km-cm	0.5	80
Pipeline Investment Subtotal			\$6,400
Offsites & Utility Investment Contingency	20%		\$2,758
Offsite & Utilities Investment			\$16,548
General Service Facilities	10%	of BLI & Utilities Investment	\$9,910
Waste Treatment	5%	of BLI Investment	\$4,130
Outside Battery Limits Investment			\$30,600
Greenfield Total Plant Investment			\$113,200
Retrofit Complexity Factor	20%	Additional Retrofit Costs	\$22,600
Total Estimated Investment			\$135,800

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Table D.6. Operating Cost Estimate: 30% Mixed Amine Solvent.

Table D.6 CO2 Removal Using 30% Mixed Amine (0.5 MEA, 0.4 MDEA, 0.1 AMP by weight), Gibson 3, Annual Estimated Operating Costs			
Plant Net Capacity		0.91 megatonnes/yr 1.01 MMshort tons/yr	
Onstream factor		0.8	
Price of Electricity		\$34.00 per MWh	
Costs in 2011 thousand US\$ unless noted			
			Costs
Plant Investment, Battery Limits (BLI)			\$99,100
Plant Investment, Outside Battery Limits (OBLI)			\$36,720
Total Fixed Capital (TFM)			\$135,820
Operating Costs, Per Year			
Variable Costs			
Losses to Energy Generation (net)			\$13,630
Materials Consumed Costs			\$4,290
Utility Costs (net)			\$3,710
Total Annual Variable Costs			\$21,630
Labor Costs			
Operating Labor (3 shifts)	6	\$62 wages/year	\$372
Maintenance Labor	3.00% of BLI		\$2,970
Control Laboratory Labor, 10% of Operating Labor	10% of Operating Labor		\$40
Direct Labor Costs			\$3,380
Maintenance Materials	3.00% of BLI		\$2,970
Operating Supplies, 12% of Operating Labor	12% of Operating Labor		\$40
Total Direct Costs			\$28,020
Plant Overhead	80% of Direct Labor Costs		\$2,700
Taxes and Insurance	1.60% of TFC		\$2,170
Cash Costs			\$32,890
Levelized Depreciation	3.3% of TFC		\$4,530
Gate Costs			\$37,420
General, Admin, Sales, Research	5% of Gate Costs		\$1,870
Production Costs, excluding Cost of Capital			\$39,300
TFC + Estimated Working Capital*			\$144,700
Annual Cost of Capital	12.5% of Capital Investment		\$18,090
Total Operating Costs			\$57,390

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<i>Cost per tonne CO2, dollars</i>	\$57.00
Additional costs per MWh dollars	\$66.0
Nominal Output of Gibson-3, MW	695 MWe
Nominal Output after existing peripheral losses, MW	635 MWe
Thermal Rating of Gibson-3, MW	1809 MWth
Share of Gibson-3 CO2 emissions processed	25.5% MW
Equivalent electricity output of share of plant emissions processed before carbon capture losses	162 MWe
Power losses from Carbon Capture and Compression, MW	38 MWe
Power output of share of plant for which emissions are processed after energy losses	124 MWe
Reduction in Power Output due to carbon capture	23.4%
Efficiency without Carbon Capture	35.1%
Efficiency with Carbon Capture	26.9%
Reduction in Efficiency	-8.22%
Annual Electrical Energy output of share of plant for which emissions are processed after carbon capture losses	869,100 MWh

* Working Capital was estimated at 25% of Annual Cash Costs plus cost of chemical and resin inventory

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Table D.7. Capital Cost Estimate: 35% Mixed Amine Solvent.

Table D.7
CO2 Removal Using 35% Mixed Amine (0.5 MEA, 0.4 MDEA, 0.1 AMP by weight), Gibson 3,
Annual Estimated Variable Costs

Costs in 2011 US\$

Onstream Factor 0.8

Losses to Energy Generation

Lost Generation From Steam Bleed to Reboiler	24.5 MWh	\$51.2 per MWh*	\$1,256	\$8,805,000
Power generated by Letdown Turbine	(4.3) MWh	\$51.2 per MWh*	\$(222)	\$(1,554,000)
Parasitic Electricity Consumption by Compressors/Pumps	15.7 MWh	\$51.2 per MWh*	\$806	\$ 5,648,000
Losses to Energy Generation			\$ 1,841	\$ 12,899,000

Materials Consumed	Number/hr	Unit Cost \$	Costs/hr	Costs/yr
Monoethanolamine (MEA)	0.11 tonnes	\$2,880 per tonne	\$302	\$2,119,000
Methyl Diethanolamine (MDEA)	0.024 tonnes	\$2,980 per tonne	\$70	\$491,000
2-Amino Methyl Propanol (AMP)	0.005 tonnes	\$4,960 per tonne	\$23	\$164,000
Stabilizer (5% of amine makeup costs)				\$139,000
Caustic Soda	0.16 tonnes	\$460 per tonne	\$74	\$516,000
Desiccant Replacement	0.0031 tonnes	\$5,000 per tonne	\$15	\$107,000
Activated Carbon	0.0024 tonnes	\$2,500 per tonne	\$6	\$43,000
Ion Exchange Resin	0.0059 tonnes	\$8,000 per tonne	\$47	\$333,000
Disposal/treatment of Selenium-containing wastes	0.1140 tonnes	\$300 per tonne	\$34	\$240,000
Other Waste Disposal	0.0650 tonnes	\$300 per tonne	\$20	\$137,000
Material Consumed Costs			\$592	\$4,289,000
Utilities				
Cooling Water	25,920 cu.m	\$0.02 per cu.m	\$497	\$3,482,000
Utility Costs			\$497	\$3,482,000
Total Variable costs			\$2,930	\$20,670,000

* 2006-2011 Five-year average bilateral contract price for Cinergy/MISO hub used as the cost of electricity

* Relative degradation rates of the individual amines were based on Ind. Eng. Chem. Res., 2009, 48 (20), pp 9061–9067:

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Table D.8. Variable Cost Estimate: 35% Mixed Amine Solvent.

Table D.8 CO2 Removal Using 35% Mixed Amine (0.5 MEA, 0.4 MDEA, 0.1 AMP by weight), Gibson 3, Estimated Fixed Capital Costs			
Plant Net Capacity, CO2 Removed/Hr			130 tonnes/hr 144 short tons/hr
Costs in 2011 thousand US\$			
Gas Cleanup and Power Plant			
Battery Limits Investment	Equipment Cost	Installation Cost	Total Cost
Absorber Columns	\$4,140	\$9,110	\$13,250
Stripper Columns	\$1,080	\$2,380	\$3,460
Heat Exchangers	\$7,600	\$11,900	\$19,500
Pressure Vessels	\$2,450	\$3,430	\$5,880
Fans	\$175	\$245	\$420
CO2 Compressors	\$6,260	\$9,350	\$15,610
Pumps	\$1,660	\$2,850	\$4,010
Let-down Turbine	\$1,500	\$1,200	\$2,700
Subtotal			\$64,830
BLI Contingency	20%	of Installed Equipment Costs	\$12,970
Battery Limits Investment			\$77,80
Tankage	Equipment Cost	Installation Cost	
Amine Solution Surge Tanks	\$260	\$310	\$570
MEA Storage Tank	\$100	\$120	\$220
MDEA Storage Tank	\$60	\$70	\$130
AMP Storage Tank	\$40	\$50	\$90
Makeup Amine Staging Tank	\$100	\$120	\$220
Makeup Amine Staging Tank	\$100	\$120	\$220
Caustic Soda Tank	\$130	\$210	\$340
Tankage Investment Subtotal			\$1,790
Utilities	Equipment Cost	Installation Cost	Investment
Cooling Water	\$3,190	\$2,200	\$5,390
Utilities Investment Subtotal			\$5,400
Pipelines from Main Plant	Unit cost, installed	Length (km)	Diameter (cm)
Flue Gas	\$16/km-cm	0.5	660
Steam	\$16/km-cm	0.5	60
Condensate	\$16/km-cm	0.5	80
Pipeline Investment Subtotal			\$6,400
Offsites & Utility Investment Contingency	20%		\$2,758
Offsite & Utilities Investment			\$16,548
General Service Facilities	10%	of BLI & Utilities Investment	\$9,910
Waste Treatment	5%	of BLI Investment	\$4,130
Outside Battery Limits Investment			\$30,600
Greenfield Total Plant Investment			\$113,200
Retrofit Complexity Factor	20%	Additional Retrofit Costs	\$22,600
Total Estimated Investment			\$135,800

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Table D.9. Operating Cost Estimate: 35% Mixed Amine Solvent

Table D.9 CO2 Removal Using 30% Mixed Amine (0.5 MEA, 0.4 MDEA, 0.1 AMP by weight), Gibson 3, Annual Estimated Operating Costs			
Plant Net Capacity		0.91 megatonnes/yr 1.01 MMshort tons/yr	
Onstream factor		0.8	
Price of Electricity		\$51.20 per MWh	
Costs in 2011 thousand US\$ unless noted			
			Costs
Plant Investment, Battery Limits (BLI)			\$93,400
Plant Investment, Outside Battery Limits (OBLI)			\$35,520
Total Fixed Capital (TFM)			\$128,920
Operating Costs, Per Year			
Variable Costs			
Losses to Energy Generation (net)			\$12,900
Materials Consumed Costs			\$4,290
Utility Costs (net)			\$3,480
Total Annual Variable Costs			\$20,670
Labor Costs			
Operating Labor (3 shifts)	6	\$62 wages/year	\$372
Maintenance Labor	3.00% of BLI		\$2,802
Control Laboratory Labor, 10% of Operating Labor	10% of Operating Labor		\$40
Direct Labor Costs			\$3,210
Maintenance Materials	3.00% of BLI		\$2,802
Operating Supplies, 12% of Operating Labor	12% of Operating Labor		\$40
Total Direct Costs			\$26,722
Plant Overhead	80% of Direct Labor Costs		\$2,570
Taxes and Insurance	1.60% of TFC		\$2,060
Cash Costs			\$31,352
Levelized Depreciation	3.3% of TFC		\$4,300
Gate Costs			\$35,650
General, Admin, Sales, Research	5% of Gate Costs		\$1,780
Production Costs, excluding Cost of Capital			\$37,400
TFC + Estimated Working Capital*			\$137,500
Annual Cost of Capital	12.5% of Capital Investment		\$17,190
Total Operating Costs			\$54,590

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<i>Cost per ton CO2, dollars</i>	<i>\$54.20</i>
<i>Cost per tonne CO2, dollars</i>	<i>\$59.80</i>
Additional costs per MWh dollars	\$61.8
Nominal Output of Gibson-3, MW	696 MWe
Nominal Output after existing peripheral losses, MW	635 MWe
Thermal Rating of Gibson-3, MW	1809 MWth
Share of Gibson-3 CO2 emissions processed	25.5% MW
Equivalent electricity output of share of plant emissions processed before carbon capture losses	162 MWe
Power losses from Carbon Capture and Compression, MW	36 MWe
Power output of share of plant for which emissions are processed after energy losses	126 MWe
Reduction in Power Output due to carbon capture	22.2%
Efficiency without Carbon Capture	35.1%
Efficiency with Carbon Capture	27.3%
Reduction in Efficiency	-7.8%
Annual Electrical Energy output of share of plant for which emissions are processed after carbon capture losses	883,400 MWh

* Working Capital was estimated at 25% of Annual Cash Costs plus cost of chemical and resin inventory

Appendix E: Discussion of Shidongkou Plant Visit

The authors visited Huaneng's Shidongkou pilot plant at Shanghai on Friday, 17 August, 2012. The plant was not operating at the time, but we discussed its operation with Huaneng engineers and plant personnel. The plant was designed to take part (4%) of the flue gas from a 600 MW coal fed power generating unit, and generate 100,000 metric tons of food grade carbon dioxide for commercial use.⁹ The installation, the world's largest carbon capture plant when it was built, cost \$24 million to build.¹⁰

Although we were only able to photograph the equipment from outside the fence, it was apparent that the compactness of the installation enabled good heat integration, where a more spread out layout would worsen the heat integration because of heat loss from long pipe runs. According to plant personnel, startup and shutdown of the carbon capture plant was fast, less than one hour. Also, the carbon capture could be run as low as 50% of the design capacity of 100,000 tonnes per year. Because of overdesign of critical pieces of equipment, Shidongkou was able, through debottlenecking, to increase the capacity to 120,000 tonnes per year.

To date, the only amine system tested at the Shidongkou location was monoethanolamine (MEA), 15-20% in water.

Figure E.1 is a photograph of process flow diagram (PFD) of the Shidongkou carbon capture plant, and Figure E.2 is a recreation of the same PFD. For comparison, Figure E.3, identical to Figure A.1, is the PFD from our Aspen model. The recreated PFD in Figure E.2 does not include the purification system to make food grade CO₂, because that level of purity is not necessary in our scenarios. In our Aspen model, the liquid CO₂ product is 99.5% pure after the five-stage compression with interstage cooling. Also, while the Shidongkou plant includes a spherical holding tank for CO₂, we have not included a CO₂ holding tank in our cost estimate, assuming that storage of CO₂ would be the responsibility of the receiver of the CO₂. Other differences exist between the PFD from Shidongkou and our Aspen model such as:

- The Shidongkou plant does not have precooling of the flue gas before the absorber.
- There is a preheater before the recycle heat exchanger (we believe this may be recovering some low-grade heat from the flue gas stream before the FGD)
- The lean solution bleed system is not shown. In this system, 10% of the lean solution is bled off, then combined with caustic soda and flashed under vacuum, the vapor to be recompressed and injected into the bottom section of the stripper.
- Shidongkou has a somewhat more rigorous process for purification of CO₂ (as the CO₂ needs to meet food-grade specifications for use in carbonated beverages)
- The stripper reboiler is not shown as a separate piece of equipment in either Figure E.2. or Figure E.3. Although the reboiler is an important component of the carbon capture equipment and the source of most of the energy usage, it is internal to the stripper, so is not shown in the figures.

Plant personnel informed us that the absorber and stripper columns were 4 meters in diameter, which is consistent with our Aspen-calculated absorber diameter of about 12 meters for the Duke system with ten times the throughput, since the diameter varies with the square root of the vapor rate. Based on this, we estimated from Figure E.4. that the column heights are approximately 20 meters.

Further photographs of equipment at the Shidongkou CO₂ capture plant are given in Figures E.5 to E.9. As noted above, it can be seen in the photographs that the system at Shidongkou is compact in area. While we did not take exact measurements, we judged the area of the CO₂ capture plant to be perhaps 200 x 50 feet in dimension.

Passivation of Carbon Steel at Gaobeidian

Huaneng passivated the system at their Beijing power plant at Gaobeidean, designed to remove 3,000 tonnes per year, using MEA and an antioxidant as the passivation agents.¹¹ This was an experiment to see if lower grades of steel than stainless could be used in CO₂ capture systems. Although they circulated the solution for 200 hours, the amine concentration fluctuated and they had to periodically adjust the concentration, so were not certain of the passivation results. Corrosion resulted in some unit operations.

Figure E.1. Photograph of Shidongkou PFD.

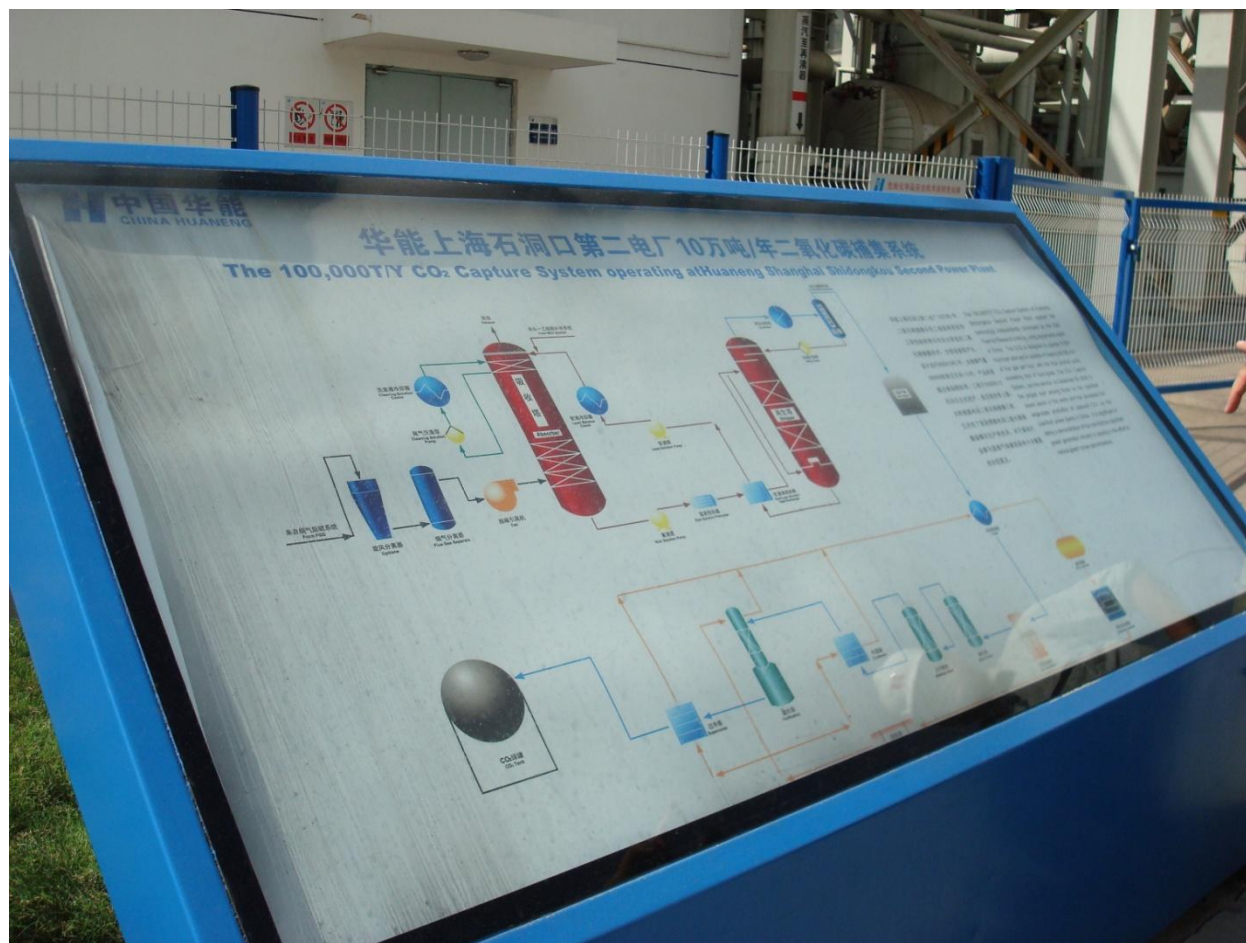


Figure E.2. Recreation of PFD for Shidongkou CO₂ Capture Plant.

The 100,000T/Y CO₂ Capture System Operating at Huaneng Shanghai Shidongkou Second Power Plant

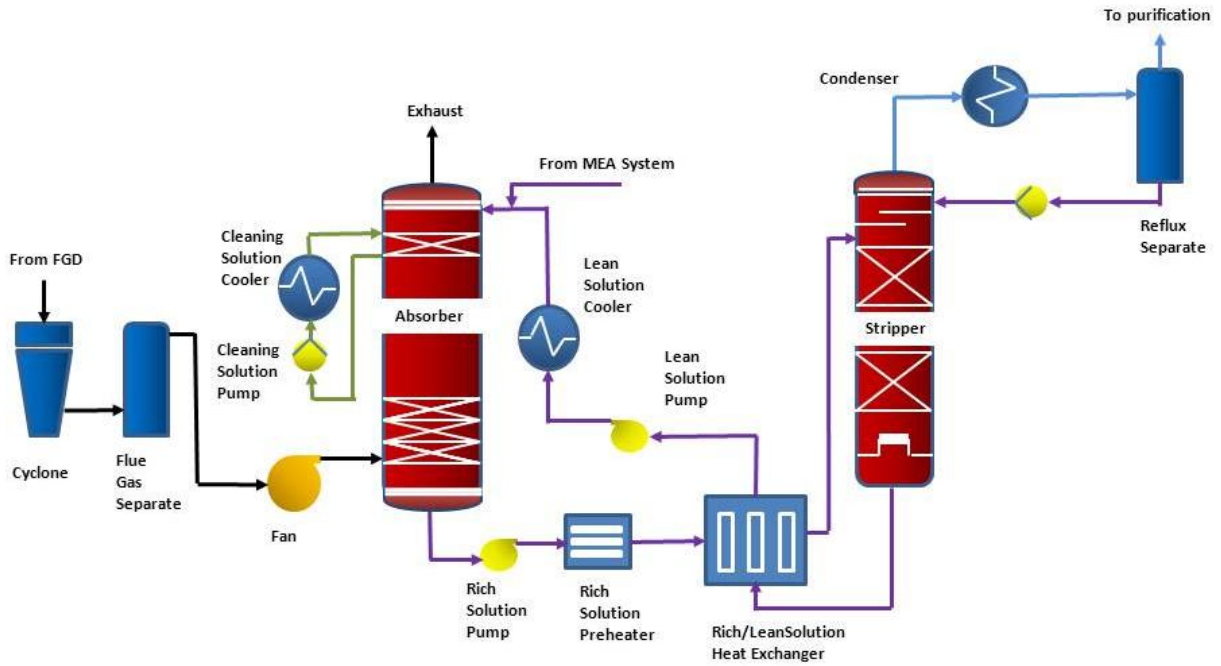


Figure E.3. Carbon Capture PFD from LLNL Aspen model.

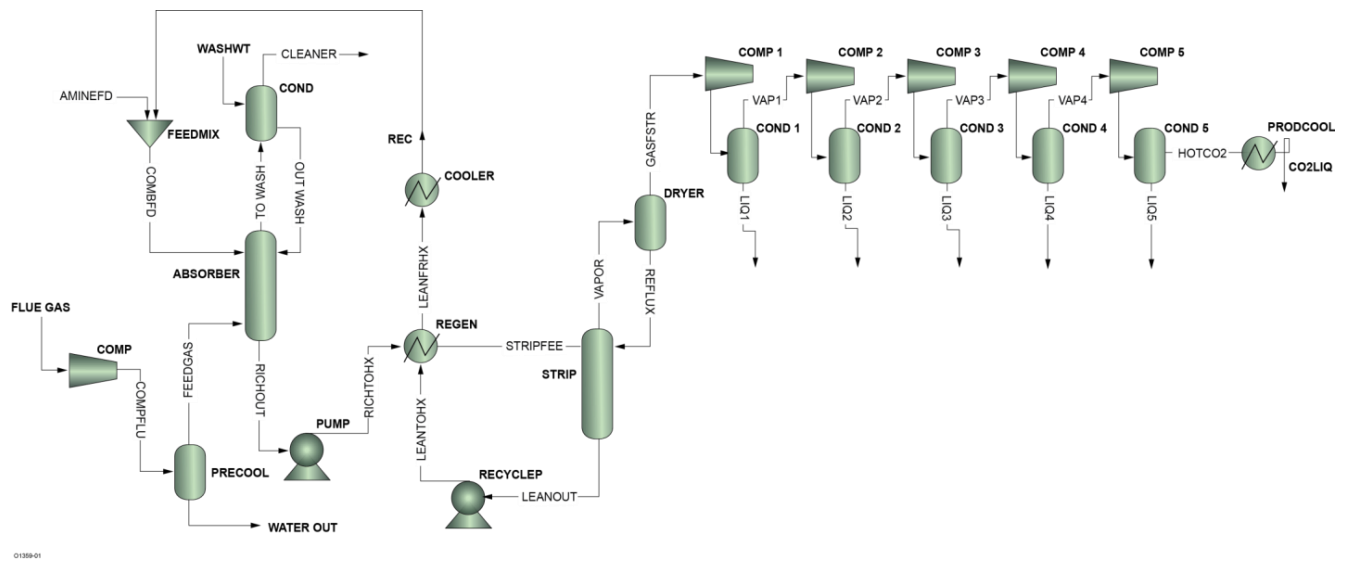


Figure E 4. Photograph Absorber and stripper columns.



Figure E.5. CO₂ Purification and Product CO₂ storage tanks.



Figure E.6. Stripper Reflux Separator.



Figure E.7. Probable Water Storage Tank.



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Figure E.8. Probable Solvent Surge Tank and Feed Amine Storage Tank.



Same floor of this platform contains the plate heat exchangers used (but are not visible from this vantage point) in the process to exchange heat between the stripper feed and stripper bottoms.

Figure E.9. Pump Sets.



Figure E.10. Amine Recovery Vacuum Flash Vessel.



This flash vessel is used to recover MEA from a purge stream from the solvent recycle. The solvent purge is adjusted to high pH using caustic soda to shift the equilibrium of the amine away from the carbamate and the amine ion form, and to un-ionized amine, which is more volatile. The now highly alkaline purged solvent is then heated and flashed in the vessel shown. MEA is volatilized and recovered; the bottoms from the flash vessel, containing heat-stable salts and heavy amine degradation products, is sent to waste treatment and disposal.

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